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INTERNATIONAL APPLICATION PUBLISH	HED U	NDER THE PATENT COOPERATION TREATY (PCT) -		
(51) International Patent Classification ⁶ :		(11) International Publication Number: WO 99/28464		
C12N 15/12, C07K 14/47, C12N 15/11, C07K 16/18, G01N 33/50	A2	(43) International Publication Date: 10 June 1999 (10.06.99)		
(21) International Application Number: PCT/US (22) International Filing Date: 27 November 1998 (2)	(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,			
(30) Priority Data: 60/066,863 28 November 1997 (28.11.9 (71) Applicant (for all designated States except US): WISI ALUMNI RESEARCH FOUNDATION [US/US];	MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).			
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(54) Title: CDNAS AND PROTEINS BELONGING TO AND METHODS OF USE	THE	 HLH-PAS SUPERFAMILY OF TRANSCRIPTION REGULATORS,		
(57) Abstract	*			
of transcription regulators. These "MOPs" (members of P.	AS) are	teins that are new and distinct members of the bHLH-PAS superfamily useful in a variety of research, diagnostic and therapeutic applications. a-inducible factors. Several other of the MOPs of the invention are		
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CDNAS AND PROTEINS INVOLVED IN HYPOXIA, CIRCADIAN AND ORPHAN SIGNAL TRANSDUCTION PATHWAYS, AND METHODS OF USE

Pursuant to 35 U.S.C. §202(c), it is acknowledged that the U.S. Government has certain rights in the invention described herein, which was made in part with funds from the National Institutes of Health, Grant Nos. P30-CA07175 and ES05703.

This application claims priority to U.S. Provisional Application Serial No. 60/066,863, filed November 28, 1997, which is incorporated by reference herein.

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FIELD OF THE INVENTION

This invention relates to the field of molecular signaling and physiological responses to external stimuli. In particular, this invention provides nucleic acid molecules and proteins that constitute new members of the bHLH-PAS superfamily of transcription regulators.

BACKGROUND OF THE INVENTION

Several publications are referenced in this application to describe the state of the art to which the invention pertains. Each of these publications is incorporated by reference herein.

The aryl hydrocarbon receptor (AH receptor or AHR), AH receptor nuclear transporter (ARNT), Drosophila single-minded gene product (SIM) and Drosophila period gene product (PER) are the founding members of an emerging superfamily of regulatory proteins. The AHR and ARNT are heterodimeric partners that transcriptionally upregulate genes involved in the metabolism of xenobiotics. The AHR is activatable by a number of widespread environmental pollutants like 2,3,7,8-

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tetrachlorodibenzo-p-dioxin (TCDD). In the absence of agonist, the AHR is primarily cytosolic and functionally repressed, presumably as the result of its tight association with Hsp90. Current models suggest that agonist binding initiates translocation of the receptor complex to the nucleus and concomitantly weakens the AHR-Hsp90 association. Within the nucleus, Hsp90 is displaced and the AHR dimerizes with its partner ARNT resulting in a bHLH-PAS heterodimer with binding specificity for DNA sequences within enhancer elements upstream of gene products that metabolize foreign chemicals. In Drosophila, SIM is master regulator of midline cell lineage in the embryonic nervous system. vitro and in vivo studies suggest that SIM may also dimerize with an ARNT-like protein to regulate enhancer sequences present in the sim, slit and Toll structural genes. The Drosophila PER protein plays a role in the maintenance of circadian rhythms. PER has been shown to form heterotypic interactions with a second Drosophila protein, TIM, in vivo, and homotypic interactions with the ARNT molecule in vitro.

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The distinguishing characteristic of these proteins is a 200-300 stretch of amino acid sequence similarity known as a PAS (PER/ARNT/SIM) domain. In the AHR, the PAS domain has been shown to encode sites for agonist binding, surfaces to support heterodimerization with other PAS domains, as well as surfaces that form tight interactions with Hsp90. In addition to the PAS domain, the AHR, ARNT and SIM also harbor a bHLH (basic helix-loop-helix) motif that plays a primary role in dimer formation. The bHLH motif is found in a variety of transcription factors that utilize homotypic interactions to regulate various aspects of cell growth and differentiation. Dimerization specificity is dictated by sequences within both the bHLH and determinants within secondary interaction surfaces, such as the "leucine zipper or PAS domains. Interestingly, these dimerization

surfaces also appear to restrict pairing to within a given bHLH protein superfamily, thus minimizing crosstalk between important cellular pathways.

Because other bHLH protein families utilize multiple homotypic interactions to provide fine control in the regulation of certain gene batteries, it is possible that additional bHLH-PAS proteins exist in the mammalian genome and that a subset of these proteins might dimerize with either the AHR or ARNT. prior to the present invention, the AHR and ARNT were the only mammalian bHLH-PAS proteins that had been identified. Accordingly, a need exists to identify and characterize other bHLH-PAS domain proteins, particularly those that are novel receptors for drugs, or are AHR or ARNT binding partners. Such molecules would find broad utility as research tools in elucidating environmentally and developmentally controlled signal transduction pathways, and also as diagnostic and therapeutic agents for detection and control of such pathways.

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SUMMARY OF THE INVENTION

This invention provides isolated nucleic acids and proteins which are new and distinct members of the bHLH-PAS superfamily of transcription regulators. These "MOPs" (members of PAS) are useful for a wide variety of research, diagnostic and therapeutic applications, as described in greater detail herein.

According to one aspect of the invention, isolated nucleic acid molecules are provided that include an open reading frame encoding a protein selected from the group consisting of: MOP2, MOP3, MOP4, MOP5, MOP6 MOP7, MOP8 and MOP9. In preferred embodiments, the open reading frame encodes a protein having an amino acid sequence substantially the same as a sequence selected from the group consisting of SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16 SEQ ID NO:17 and SEQ ID NO:18. The nucleic acid

molecules of the invention preferably comprise sequences substantially the same as a sequence selected from the group consisting of SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 SEQ ID NO:8 and SEQ ID NO:9.

According to another aspect of the invention, isolated MOP proteins are provided, which are products of expression of part or all of the open reading frames of the aforementioned nucleic acid molecules.

According to another aspect of the invention, recombinant DNA molecules are provided, which comprise MOP encoding nucleic acid molecules, operably linked to vectors for transforming cells. Cells transformed with those recombinant DNA molecules are also provided, as well as cellular assay systems utilizing those recombinant molecules.

According to another aspect of the invention, oligonucleotides between about 10 and about 100 nucleotides in length are provided, which specifically hybridize with portions of the MOP-encoding nucleic acid molecules.

According to another aspect of the invention, antibodies are provided which are immunologically specific for part or all of any of the MOP2-MOP8 proteins of the invention.

According to another aspect of the invention, assays and other methods of using the aforementioned MOP nucleic acids, proteins and immunospecific antibodies are provided.

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BRIEF DESCRIPTION OF THE DRAWINGS

FIGURE 1. Schematic representation of a generic bHLH-PAS member and the corresponding region where EST "hits" occurred. Top, schematic of a generic bHLH-PAS family member. The hatched box represents the bHLH region, the overlined area represents the PAS domain with the characteristic "A" and "B" repeats in white.

The variable C terminus is boxed in white. A bold line representing the region in a generic bHLH-PAS member where the homology occurs is indicated next to the original Gen BankTM accession number for each identified EST (MOP1=T10821, MOP2 = T70415, MOP3 = T77200 and F06906, MOP4 = R58054, MOP5 = R67292; see Table 1).

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alignment of the PAS domains. The amino acid sequence including a CLUSTAL alignment of the bHLH-PAS domains is depicted. The CLUSTAL alignment was performed using the MEGALIGN program (DNASTAR, Madison, WI) with a PAM250 weight table using the following parameters: Ktuple = 1, Gap Penalty = 3, Window = 5. Amino acid boundaries for the residues encompassing the bHLH and PAS domains of the MOPs were defined based on previous observations. The bHLH domain is boxed, while the basic region is specified by a vertical line. The PAS domain is underlined, while the "A" and "B" repeats of the PAS domain are boxed. Consensus (60%) residues in the PAS domain are denoted with an asterisk.

FIGURE 3. Yeast two-hybrid analysis. In vivo interaction of MOPs with dioxin signaling pathway. Fig. 3A, schematic of AHR, ARNT, and LexA fusion constructs. Panel shows a schematic of the AHR, with the PAS domain (black) with the characteristic "A" and "B" repeats (white), the bHLH domain (striped), and the variable C terminus (white). The transcriptionally active glutamine rich domain is indicated with "Q" (shaded box). LexA fusion proteins are indicated with the N terminus of LexA DNA-binding protein fused to bHLH-PAS domains of MOPs 1-4, and ARNT. The LexAAHR construct contains the bHLH-PAS domains and the C terminus minus the transcriptionally active Q-rich region (see Example 1, "Materials and Methods"). Fig. 3B, relative interaction of LexA fusion proteins with the AHR or ARNT. Galacto-Light assays were performed on yeast extracts prepared from colonies expressing LexAMOPs, LexAAHR, or LexAARNT and ARNT or AHR

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in the presence and absence of 1 μ M β NF. Assays were performed in triplicate, and then the relative light units normalized to the LexAAHR-ARNT + β NF condition internally (set as 100%). The stippled bars represent LexA fusion proteins co-expressed with the full-length AHR in the presence of 1 μ M β NF, the striped bars represent the fusion proteins co-expressed with the full length AHR in the absence of ligand, the shaded bars indicate the fusion proteins co-expressed with the full-length ARNT, and the open bar indicates LexAAHR co-expressed with full-length ARNT in the presence of 1 μ M β NF.

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FIGURE 4. Schematic comparison of homology of PAS family members. A dendrogram was prepared from the 15 primary amino acid CLUSTAL alignment above using the MEGALIGN program. The CLUSTAL alignment was performed using the MEGALIGN program (DNASTAR, Madison, WI) with PAM250 weight table using the following parameters: Ktuple =1, Gap Penalty = 3, Window = 5. Amino acid 20 boundaries for the residues encompassing the bHLH and PAS domains of the MOPs were defined based on previous observations. The amino acid boundaries are as follows: huMOP1/HIF1 (91-342), huMOP2 (90-342), huMOP3 (148-439), huMOP4 (87-350), huMOP5 (32-296), huAHR (117-385), huARNT 25 (167-464), drSIM (82-356), drPer (232-496) bsKINA (27-248), huSRC-1(115-365), muARNT2 (141-437, muSIM1 (83-331), muSIM2 (83-332), drSIMILAR (174-419), and drTRH (145-471). The scale at the bottom indicates number of amino acid residue substitutions. PAS family members 30 that interact with HSP90, interact with the AHR, and interact with ARNT by the coimmunoprecipitation method are indicated by a +, whereas members that do not interact are indicated by a -. An α denotes a bHLH-PAS member whose cDNA is not complete. Note that these 35 interactions occur in vitro and may or may not be physiologically relevant. Where appropriate, the reference is included in parentheses.

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FIGURE 5. Interaction panel of LexAbHLH-PAS fusion proteins with full-length MOP3 and ARNT. Fig. 5A: Schematic representation of the LexAbHLHPAS "bait" and the full-length "fish." The bHLH and PAS domains are 5 boxed. The "A" and "B" repeats of the PAS domains are indicated. The transactivation domain of the full-length "fish" is indicated. Fig. 5B: LexA fusion protein plasmids containing the bHLH-PAS domains of $HIFl\alpha$, $HIF2\alpha$, MOP3, MOP4, AHR, ARNT, and CLOCK were coexpressed with 10 plasmids harboring full-length MOP3 and ARNT (see Materials and Methods). LexAAHR interactions were assayed on plates containing 1 μ M β -naphthoflavone. After incubation, an 5-bromo-4-chloro-3-indolyl 13- β galactoside overlay assay was performed. ++, A strong 15 interaction, turning blue within 2 hr; +, a weaker interaction, turning blue between 8 hr and overnight; and -, a negative interaction after overnight incubation. The experiment was performed three times with identical results.

20 FIGURE 6. The consensus DNA binding site for MOP3-MOP4 heterodimer in vitro. Ten selected DNA sequences bound by the MOP3-MOP4 complex are indicated with the E-box core boxed. Underneath, the M34 consensus is indicated. Nucleotide positions relative to the E-box core are shown. Bases in uppercase are randomer derived, 25 while bases in lower case are primer derived.

FIGURE 7. Cloning of MOP7. The positions of the original EST clone (AA028416) and RACE products are shown as dark lines with the mMOP7 ORF shown as an open The PCR primers used are posted below the corresponding fragments and the plasmid numbers are marked on the side. The GenBank Accession Number for mouse MOP7 cDNA is AF060194.

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FIGURE 8. The splicing site within mouse MOP7 35 ORF are compared with those previously reported for mHIF1 α and hHIF2 α . The numbers of amino acids at which the splicing occurs are marked underneath the sequence.

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The conserved splicing sites are defined as the splicing sites of $HIF1\alpha$ and $HIF2\alpha$ that are within one amino acid of the corresponding MOP7 splicing site on the aligned sequence map using CLUSTAL method. These sites are marked with lines between different ORFs (see GenBank Accession Numbers AF079140-079153 for detailed sequences of mMOP7 splice sites.

DETAILED DESCRIPTION OF THE INVENTION

10 I. <u>Definitions</u>

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Various terms relating to the biological molecules of the present invention are used hereinabove and also throughout the specifications and claims. The terms "substantially the same," "percent similarity" and "percent identity" are defined in detail below.

With reference to nucleic acids of the invention, the term "isolated nucleic acid" is sometimes used. This term, when applied to DNA, refers to a DNA molecule that is separated from sequences with which it is immediately contiguous (in the 5' and 3' directions) in the naturally occurring genome of the organism from which it was derived. For example, the "isolated nucleic acid" may comprise a DNA molecule inserted into a vector, such as a plasmid or virus vector, or integrated into the genomic DNA of a procaryote or eucaryote. An "isolated nucleic acid molecule" may also comprise a cDNA molecule.

With respect to RNA molecules of the invention, the term "isolated nucleic acid" primarily refers to an RNA molecule encoded by an isolated DNA molecule as defined above. Alternatively, the term may refer to an RNA molecule that has been sufficiently separated from RNA molecules with which it would be associated in its natural state (i.e., in cells or tissues), such that it exists in a "substantially pure" form (the term "substantially pure" is defined below).

With respect to protein, the term "isolated protein" or "isolated and purified protein" is sometimes

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used herein. This term refers primarily to a protein produced by expression of an isolated nucleic acid molecule of the invention. Alternatively, this term may refer to a protein which has been sufficiently separated from other proteins with which it would naturally be associated, so as to exist in "substantially pure" form.

The term "substantially pure" refers to a preparation comprising at least 50-60% by weight the compound of interest (e.g., nucleic acid, oligonucleotide, protein, etc.). More preferably, the preparation comprises at least 75% by weight, and most preferably 90-99% by weight, the compound of interest. Purity is measured by methods appropriate for the compound of interest (e.g. chromatographic methods, agarose or polyacrylamide gel electrophoresis, HPLC analysis, and the like).

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With respect to antibodies of the invention, the term "immunologically specific" refers to antibodies that bind to one or more epitopes of a protein of interest, but which do not substantially recognize and bind other molecules in a sample containing a mixed population of antigenic biological molecules.

With respect to single stranded nucleic acids, particularly oligonucleotides, the term "specifically hybridizing" refers to the association between two single-stranded nucleotide molecules of sufficiently complementary sequence to permit such hybridization under pre-determined conditions generally used in the art (sometimes termed "substantially complementary"). In particular, the term refers to hybridization of an oligonucleotide with a substantially complementary sequence contained within a single-stranded DNA or RNA molecule of the invention, to the substantial exclusion of hybridization of the oligonucleotide with single-stranded nucleic acids of non-complementary sequence.

The term "promoter region" refers to the transcriptional regulatory regions of a gene. In the

present invention, the use of SV40, TK, Albumin, SP6, T7 gene promoters, among others, is contemplated.

The term "selectable marker gene" refers to a gene encoding a product that, when expressed, confers a selectable phenotype such as antibiotic resistance on a transformed cell.

The term "reporter gene" refers to a gene that encodes a product which is easily detectable by standard methods, either directly or indirectly.

The term "operably linked" means that the regulatory sequences necessary for expression of the coding sequence are placed in the DNA molecule in the appropriate positions relative to the coding sequence so as to enable expression of the coding sequence. This same definition is sometimes applied to the arrangement of transcription units and other transcription control elements (e.g. enhancers) in an expression vector.

20 II. Characterization of MOPS 1-9

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Our hypothesis in accordance with the present invention was that additional bHLH-PAS proteins are encoded in the mammalian genome and that some of these proteins are involved in mediating the pleiotropic response to potent AHR agonists like TCDD. It has been observed that other bHLH superfamilies employ multiple dimeric partnerships to control complex biological processes, such as myogenesis (MyoD/myogenin), cellular proliferation (Myc, Max, Mad) and neurogenesis (achaete-scute/daughterless). The observation that bHLH proteins often restrict their dimerization to within members of the same gene family (i.e., "homotypic interactions") and that this restriction may occur as the result of constraints imposed by both primary (e.g., bHLH) and secondary dimerization surfaces (e.g., leucine zippers and PAS), prompted us to screen for additional bHLH-PAS proteins and test each protein for its capacity to interact with either the AHR or ARNT. The ultimate

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objective was to identify MOPs that were physiologically relevant partners of either the AHR or ARNT in vivo. Our prediction was that such proteins might respond to or modulate the AHR signaling pathway or other signaling pathways involving ARNT.

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To rapidly identify expressed genes, the "expressed sequence tag" (EST) approach was developed, whereby a cDNA library is constructed and randomly selected clones are sequenced from both vector arms (Adams et al., Science 252: 1651-1656, 1991). These partial sequences, generally 200-400 bp, are deposited in a number of computer databases that can be readily analyzed using a variety of search algorithms. 1996, the I.M.A.G.E. Consortium has deposited over 300,000 human ESTs, generated from different tissues and developmental time periods into publicly accessible databases, identifying approximately 40,000 unique cDNA clones (Lennon et al., Genomics 33: 151-152, 1996). availability of these sequences and plasmids harboring their corresponding cDNA clones provided a means by which to identify novel members of the bHLH-PAS family by nucleotide homology screening of available EST databases.

At the time this invention was initiated, the human AHR and ARNT and the drosophila SIM and PER were the only PAS protein that had been described. Therefore, we used the nucleotide sequences encoding their PAS domains as query sequences in BLASTN searches of the available EST databases. Using this strategy in an iterative fashion and confirming each hit with a reverse BLASTX search, we have identified eight cDNAs referred to herein as members of the PAS superfamily, or "MOPs". Using PCR, we were able to obtain the complete ORFs of MOPs 1-4, and extensive but incomplete ORFs of MOP5. We have also identified four more MOPs, MOPs 6, 7, 8 and 9, and obtained their complete ORFs.

While MOPs 1-5 were being characterized, Wang and colleagues identified two factors involved in

cellular response to hypoxia, HIF1α and HIF1β. These proteins are identical to MOP1 and ARNT, respectively (Wang et al., Proc. Natl. Acad. Sci. USA 92: 5510-5514, 1995). Thus, of the nine MOPs we have cloned, seven have not been previously characterized. For consistency herein, we describe MOP1 extensively, and describe heretofore undisclosed methods of using MOP1.

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The experimental approach taken in accordance with the present invention has significantly expanded the 10 number of known members of the emerging bHLH-PAS superfamily of transcriptional regulators. Along with the MOPs described herein, five additional mammalian bHLH-PAS proteins have been identified, $HIFl\alpha$ (MOP1, as described above), SIM1, SIM2, ARNT2, and SRC-1 (Wang et al., 1995, supra; Hirose et al., Mol. Cell. Biol. 16: 15 1706-1713, 1996; Fan et al., Mol. Cell. Neurosci. 7: 1-16, 1996; Ema et al., Mol. Cell. Biol. <u>16</u>: 5865-5875, 1996; Chen et al., Nat. Genet. <u>10</u>: 9-10, 1995; and Kamei et al., Cell 85: 403-414, 1996). To compare amino acid sequences of these proteins, we performed a CLUSTAL 20 alignment with the bHLH-PAS domains of MOPs 1-5 and all the known family members using a PAM250 residue weight table (Higgins & Sharp, Gene (Amst.) 73: 237-244, 1988). The two most related members were MOP1/HIF1 α and MOP2, 25 which shared 66% identity in the PAS domain. A comparison of these two proteins reveals only a single amino acid difference in the basic region and 83% identity in the HLH region. This sequence similarity is in agreement with our contention (discussed in Example 1) that $MOP1/HIF1\alpha$ and MOP2 function analogously, 30 interacting with the same heterodimeric partners and binding similar enhancer sequences in vivo. A comparison of MOP3 and ARNT and a comparison of MOP5 and SIM reveal 40% and 38% identity in the PAS domain, respectively. The basic regions of MOP3 and ARNT have only three 35 substitutions, while the HLH domains share 66% identity,

again suggesting that the two proteins may regulate

similar or identical enhancer sequences (half sites).

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A CLUSTAL alignment of the C-termini of MOPs 1-5 and the previously identified PAS members demonstrated that these regions are not well conserved (data not shown) (Burbach et al., Proc. Natl. Acad. Sci. USA 89: 8185-8189, 1992). This lack of conservation may indicate that the C-termini of these genes have divergent functions, or that the functions harbored in the C-termini can be accomplished by a variety of different sequences. For example, the C-termini of the AHR, ARNT, and SIM all harbor potent transactivation domains, yet display little sequence homology.

To characterize the evolutionary and functional relationships of these proteins, we performed a parsimony analysis to identify functionally related subsets. 15 dendrogram representing the primary amino acid relationship between the PAS domains of these proteins is illustrated in Figure 4. This schematic suggests that major groups exist for eukaryotic PAS family members. The AHR, drSIMILAR, MOP1/HIF1\alpha, MOP2, drTRACHEALESS, 20 MOP5, and SIM exist in one group, ARNT, muARNT2, MOP3, and MOP4 in another and PER and huSRC-1 exist in their own groups. Interestingly, this pattern reflects what is known functionally about the existing PAS members. AHR, SIM, MOP1/HIF1 α and MOP2 have all been shown to 25 heterodimerize with the ARNT molecule and bind DNA. Additionally, the AHR and SIM are known to interact with HSP90, a chaperonin protein necessary for the signaling of the AHR and a number of steroid receptor family members in response to ligand. Based on these groupings, 30 MOP5 may also be an ARNT-interacting protein and a candidate for interacting with Hsp90 and being activated by small molecule ligands. The observation that ARNT has been shown to be capable of forming DNA binding homodimers and as heterodimers with a number of 35 previously identified members of the bHLH-PAS family (at least in vitro), suggests that it plays a role in a

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number of biological processes. Based on their similarity with ARNT, MOP3 and MOP4 may be candidates for binding DNA as homodimers, or for interacting with multiple bHLH-PAS members, possibly from the AHR group.

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In addition to the relevance of the above data to TCDD signaling, they also reveal additional factors important to cellular responses to hypoxic stress.

HIF1\(\alpha\)/MOP1 and MOP2 appear to share a common dimeric partner - ARNT, and are capable of regulating a common battery of genes. This notion is supported by three lines of evidence: (1) both MOP1 and MOP2 interact with ARNT as defined by coimmunoprecipitation or two-hybrid assay; (2) they have similar DNA half-site specificities when complexed with ARNT; and (3) they are both transcriptionally active from TACGTG enhancers in vivo. The observation that HIF1\(\alpha\)/MOP1 and MOP2 have markedly different tissue distributions suggests that these two proteins may be regulating similar batteries of genes in response to different environmental stimuli.

Alternatively, these proteins may be involved in restricting expression of certain groups of genes regulated by TACGTG-dependent enhancers. Finally, it is now known that MOP2 and MOP7 are subunits of a "HIF1-like" complex (i.e. a "HIF2α" and a "HIF3α, respectively) that regulates hypoxia responsive genes in distinct sets of tissues.

From the foregoing discussion, it can be seen that, while the MOPs share certain common features among themselves and with other new members of the bHLH-PAS superfamily, each of MOPs 2 - 9 is a distinctive and unique member of that family. cDNA and deduced amino acid sequences for each of MOPs 1 - 9 is set forth at the end of this specification. General features of each MOP are summarized below. In addition, MOPs 1-5 are described in great detail in Example 1, MOP3 is specifically described in Example 2 and MOP 7 is described in Example 3.

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MOP1: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP1 are set forth herein as SEQ ID NOS: 1 and 10, respectively. The cDNA includes a complete coding sequence for MOP1. As discussed above, MOP1 is known more commonly in the literature as HIF (Hypoxia-Inducible Factor)-1α (Wang et al., 1995, supra). The factor is induced by low oxygen. It interacts with HSP90 and with ARNT (AHR's binding partner). The ARNT-dimerized factor regulates expression of erythropoietin, among other genes.

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MOP2: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP2 are set forth herein as SEQ ID NOS: 2 and 11, respectively. The cDNA includes a complete coding sequence for MOP2. MOP2 appears to be related structurally and functionally to MOP1. Similar to MOP1, MOP2 interacts with ARNT, but not AHR, and drives transcription in its ARNT-dimerized form. Unlike MOP1, MOP2 does not appear to interact significantly with HSP90. MOP2 is induced by low oxygen and may be involved in hypoxia responses in different cells and tissues than is MOP1. MOP2 is sometimes referred to herein as $\rm HIF2\alpha$.

MOP3: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP3 are set forth herein as SEQ ID NOS: 3 and 12, respectively. The cDNA includes a complete coding sequence for MOP3. MOP 3 and MOP 4 are related to each other as binding partners, analogous to ARNT and AHR, respectively. As described in greater detail in Example 2, in addition to being a specific partner for MOP4, MOP3 is a general dimerization partner for a subset of the bHLH/PAS superfamily of transcriptional regulators. MOP3 interacts with MOP4, CLOCK, HIF1 α and HIF2 α . The MOP3-MOP4 heterodimer binds a CACGTGA-containing DNA element. Moreover, MOP3-MOP4 and MOP3-CLOCK complexes bind this element in COS-1 cells and drive transcription from a linked luciferase reporter gene. A high-affinity DNA binding site has also been deduced for a MOP3-HIF1 α complex (TACGTGA). MOP3-HIF1 α

and MOP3-HIF2 α heterodimers bind this element, drive transcription, and respond to cellular hypoxia.

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MOP3 also binds HSP90, and may be conditionally activated (like AHR) depending on whether it is bound to HSP90 (see Example 1) (of the MOP3/MOP4 dimerization pair, one appears to be conditionally activated, but as yet it is unclear which one). Evidence from Drosophila and rat suggest that MOP3 (cycle/bMAL1b) is regulated in a circadian manner.

MOP3 expression appears to be controlled by alternate 5' promoter regions. MOP3 mRNA expression overlaps in a number of tissues with each of its four potential partner molecules in vivo.

MOP4: The nucleotide and deduced amino acid

sequences of a cDNA encoding MOP4 are set forth herein as

SEQ ID NOS: 4 and 13, respectively. The cDNA includes an
apparently complete coding sequence for MOP4. MOP4
appears to be a human ortholog of a recently identified
murine gene called "Clock", for its involvement in

circadian rhythms (King et al., Cell 89: 641-653). MOP4
also interacts with HSP90 and, as discussed above, is the
dimerization partner of MOP3, and may be conditionally
activated. MOP4 appears to be localized in the
cytoplasm.

MOP5: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP5 are set forth herein as SEQ ID NOS: 5 and 14, respectively. The cDNA includes a partial coding sequence for MOP5; however a complete coding sequence for MOP5 has become publicly available subsequent to the making of the present invention (GenBank Accession No. U77968, submitted 11/11/96, published 1/21/97 by Zhou et al., Proc. Natl. Acad. Sci. USA 94: 713-718).

MOP6: The nucleotide and deduced amino acid

sequences of a cDNA encoding MOP6 (of human origin) are
set forth herein as SEQ ID NOS: 6 and 15, respectively.

The cDNA includes a complete coding sequence for MOP6.

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The nucleotide sequence of MOP6 is fairly unique. It is most similar in the 5' region to the bHLH-PAS member trachealess, which suggests that MOP6 may be a regulator (developmental or otherwise) of hypoxia. Functional data shows that MOP6 forms a partnership with ARNT and drives a hypoxia responsive element.

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MOP7: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP7 are set forth herein as SEQ ID NOS: 7 and 16, respectively. The cDNA includes a complete coding sequence for MOP7. In accordance with 10 this invention, MOP7 has been characterized as a new hypoxia-inducible factor, and therefore is sometimes referred to herein as HIF3α. The expression profile of MOP7 is as follows: testis, thymus > [lung, brain, heart, liver, skeletal muscle] > [skin, stomach, small 15 intestine, kidney]. This expression profile is distinct from any of MOP1, MOP2, MOP3, AHR and ARNT, suggesting a different functional role for MOP7. MOP7 is most closely related to MOP1/HIF1 α and MOP2 (HIF2 α), as described in 20 greater detail in Example 3. Accordingly, MOP7 is likely to regulate the same genes as does HIF1 α and HIF2 α , as evidenced by its dimerization with the same partners (ARNT, MOP3) and recognition of the same core response element. This, combined with the unique tissue-specific 25 expression of MOP7 suggests that it may have a functional role associated with response to low oxygen in the tissues in which it is expressed.

MOP8: The nucleotide and deduced amino acid sequences of a cDNA encoding MOP8 are set forth herein as SEQ ID NOS: 8 and 17, respectively. The cDNA includes a complete coding sequence for MOP8. Like MOP4 and MOP3, MOP8 may be involved in regulation of circadian rhythm. MOP8 shows sequence similarity to other genes involved in the circadian pathway (human PER, Drosophila PER, human RIGUI).

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MOP9: The nucleotide and deduced amino acid sequence of a cDNA encoding MOP9 are set forth herein as SEQ ID NOS: 9 and 18, respectively. Two ESTs (GenBank AA577389, AA576971) corresponding to a novel bHLH-PAS protein homologous to MOP3/bMAL1 were identified by TBLASTN searches of the Drosophila homolog of MOP3. Upon characterization, these clones were revealed to be truncated, and one of which appeared to be a splice variant. The cDNA was cloned from human brain mRNA, and alternative 5' splicing was found probably reflecting multiple promoters. A BLASTX search of the MOP 9 sequence reveals that it displays extended homology to MOP3 (E-154). These data suggest that MOP9 also pairs with CLOCK and MOP4 and binds an E-box element with flanking region specificity.

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Although specific MOP clones are described and exemplified herein, this invention is intended to encompass nucleic acid sequences and proteins from humans 20 and other species that are sufficiently similar to be used interchangeably with the exemplified MOP nucleic acids and proteins for the purposes described below. will be appreciated by those skilled in the art that MOPencoding nucleic acids from diverse species, and particularly mammalian species, should possess a 25 sufficient degree of homology with human MOPs so as to be interchangeably useful in various applications. present invention, therefore, is drawn to MOP-encoding nucleic acids and encoded proteins from any species in 30 which they are found, preferably to MOPs of mammalian origin, and most preferably to MOPs of human origin. Additionally, in the same manner that structural homologs of human MOPs are considered to be within the scope of this invention, functional homologs are also considered 35 to be within the scope of this invention.

Allelic variants and natural mutants of SEQ ID NOS: 1-9 or 10-17 are likely to exist within the human

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genome and within the genomes of other species. Because such variants are expected to possess certain differences in nucleotide and amino acid sequence, this invention provides isolated MOP-encoding nucleic acid molecules having at least about 65% (and preferably over 75%) sequence homology in the coding region with the nucleotide sequences set forth as SEQ ID NOS: 1-9 (and, most preferably, specifically comprising the coding regions of any of SEQ ID NOS: 1-9). This invention also provides isolated MOPs having at least about 75% (preferably 85% or greater) sequence homology with the amino acid sequence of SEQ ID NOS: 10-18. Because of the natural sequence variation likely to exist among the MOPs and nucleic acids encoding them, one skilled in the art would expect to find up to about 25-35% nucleotide sequence variation, while still maintaining the unique properties of the MOPs of the present invention. Such an expectation is due in part to the degeneracy of the genetic code, as well as to the known evolutionary success of conservative amino acid sequence variations, which do not appreciably alter the nature of the protein. Accordingly, such variants are considered substantially the same as one another and are included within the scope of the present invention.

"substantially the same" refers to nucleic acid or amino acid sequences having sequence variation that do not materially affect the nature of the protein. With particular reference to nucleic acid sequences, the term "substantially the same" is intended to refer to the coding region and to conserved sequences governing expression, and refers primarily to degenerate codons encoding the same amino acid, or alternate codons encoding conservative substitute amino acids in the encoded polypeptide. With reference to amino acid sequences, the term "substantially the same" refers generally to conservative substitutions and/or variations

in regions of the polypeptide not involved in determination of structure or function of the protein. The terms "percent identity" and "percent similarity" are also used herein in comparisons among amino acid sequences. These terms are intended to be defined as they are in the UWGCG sequence analysis program (Devereaux et al., Nucl. Acids Res. 12: 387-397, 1984), available from the University of Wisconsin.

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The following description sets forth the general procedures involved in practicing the present invention. To the extent that specific materials are mentioned, it is merely for purposes of illustration and is not intended to limit the invention. Unless otherwise specified, general cloning procedures, such as those set forth in Sambrook et al., Molecular Cloning, Cold Spring Harbor Laboratory (1989) (hereinafter "Sambrook et al.") or Ausubel et al. (eds) Current Protocols in Molecular Biology, John Wiley & Sons (1998) (hereinafter "Ausubel et al.") are used.

III. Preparation of MOP Nucleic Acid Molecules, MOP Proteins and Anti-MOP Antibodies

A. <u>Nucleic Acid Molecules</u>

Nucleic acid molecules encoding the MOPs of the invention may be prepared by two general methods: (1) They may be synthesized from appropriate nucleotide triphosphates, or (2) they may be isolated from biological sources. Both methods utilize protocols well known in the art.

The availability of nucleotide sequence information, such as a full length cDNA having any of SEQ ID NOS: 1-9, enables preparation of an isolated nucleic acid molecule of the invention by oligonucleotide synthesis. Synthetic oligonucleotides may be prepared by the phosphoramadite method employed in the Applied Biosystems 38A DNA Synthesizer or similar devices. The resultant construct may be purified according to methods known in the art, such as high performance liquid

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chromatography (HPLC). Long, double-stranded polynucleotides, such as a DNA molecule of the present invention, must be synthesized in stages, due to the size limitations inherent in current oligonucleotide synthetic methods. Thus, for example, a several-kilobase double-stranded molecule may be synthesized as several smaller segments of appropriate complementarity. Complementary segments thus produced may be annealed such that each segment possesses appropriate cohesive termini for attachment of an adjacent segment. Adjacent segments may be ligated by annealing cohesive termini in the presence of DNA ligase to construct an entire double-stranded molecule. A synthetic DNA molecule so constructed may then be cloned and amplified in an appropriate vector.

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Nucleic acid sequences encoding MOPs may be isolated from appropriate biological sources using methods known in the art. In a preferred embodiment, cDNA clones are isolated from libraries of human origin. In an alternative embodiment, genomic clones encoding MOPs may be isolated. Alternatively, cDNA or genomic clones encoding MOPs from other species, preferably mammalian species, may be obtained.

In accordance with the present invention, nucleic acids having the appropriate level sequence homology with the coding regions of any of Sequence I.D. Nos. 1-9 may be identified by using hybridization and washing conditions of appropriate stringency. For example, hybridizations may be performed, according to the method of Sambrook et al., using a hybridization solution comprising: 5X SSC, 5X Denhardt's reagent, 1.0% SDS, 100 µg/ml denatured, fragmented salmon sperm DNA, 0.05% sodium pyrophosphate and up to 50% formamide. Hybridization is carried out at 37-42°C for at least six hours. Following hybridization, filters are washed as follows: (1) 5 minutes at room temperature in 2X SSC and 1% SDS; (2) 15 minutes at room temperature in 2X SSC and 0.1% SDS; (3) 30 minutes-1 hour at 37°C in 1X SSC and 1% SDS; (4) 2

hours at 42-650in 1X SSC and 1% SDS, changing the solution every 30 minutes.

One common formula for calculating the stringency conditions required to achieve hybridization between nucleic acid molecules of a specified sequence homology (Sambrook et al., 1989):

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 $T_m = 81.5$ °C + 16.6Log [Na+] + 0.41(% G+C) - 0.63 (% formamide) - 600/#bp in duplex

As an illustration of the above formula, using [N+] = [0.368] and 50% formamide, with GC content of 42% and an average probe size of 200 bases, the T_m is 57°C. The T_m of a DNA duplex decreases by 1 - 1.5°C with every 1% decrease in homology. Thus, targets with greater than about 75% sequence identity would be observed using a hybridization temperature of 42°C.

Nucleic acids of the present invention may be maintained as DNA in any convenient cloning vector. In a preferred embodiment, clones are maintained in plasmid cloning/expression vector, such as pGEM-T (Promega Biotech, Madison, WI) or pBluescript (Stratagene, La Jolla, CA), either of which is propagated in a suitable E. coli host cell.

MOP nucleic acid molecules of the invention 25 include cDNA, genomic DNA, RNA, and fragments thereof which may be single- or double-stranded. Thus, this invention provides oligonucleotides (sense or antisense strands of DNA or RNA) having sequences capable of hybridizing with at least one sequence of a nucleic acid 30 molecule of the present invention, such as selected segments of the cDNA having any of SEQ ID NOS: 1-9. oligonucleotides are useful as probes for detecting MOP genes or mRNA in test samples of cells, tissue or other biological sources, e.g. by PCR amplification, or for the 35 positive or negative regulation of expression of MOP genes at or before translation of the mRNA into proteins.

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B. Proteins

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MOP proteins of the present invention may be prepared in a variety of ways, according to known methods. The proteins may be purified from appropriate sources, e.g., cultured or intact cells or tissues.

Alternatively, the availability of nucleic acids molecules encoding MOPs enables production of the MOP proteins using in vitro expression methods known in the art. For example, a cDNA or gene may be cloned into an appropriate in vitro transcription vector, such a pSP64 or pSP65 for in vitro transcription, followed by cell-free translation in a suitable cell-free translation system, such as wheat germ or rabbit reticulocytes. In vitro transcription and translation systems are commercially available, e.g., from Promega Biotech, Madison, Wisconsin or BRL, Rockville, Maryland.

According to a preferred embodiment, larger quantities of MOP proteins may be produced by expression in a suitable procaryotic or eucaryotic system. For example, part or all of a DNA molecule, such as any of the cDNAs having SEQ ID NOS: 1-9, may be inserted into a plasmid vector adapted for expression in a bacterial cell (such as E. coli) or a yeast cell (such as Saccharomyces cerevisiae), or into a baculovirus vector for expression in an insect cell. Such vectors comprise the regulatory elements necessary for expression of the DNA in the host cell, positioned in such a manner as to permit expression of the DNA in the host cell. Such regulatory elements required for expression include promoter sequences, transcription initiation sequences and, optionally, enhancer sequences.

The MOPs produced by gene expression in a recombinant procaryotic or eucyarotic system may be purified according to methods known in the art. In a preferred embodiment, a commercially available expression/secretion system can be used, whereby the recombinant protein is expressed and thereafter secreted

from the host cell, to be easily purified from the surrounding medium. If expression/secretion vectors are not used, an alternative approach involves purifying the recombinant protein by affinity separation, such as by immunological interaction with antibodies that bind specifically to the recombinant protein. Such methods are commonly used by skilled practitioners. The MOP proteins of the invention, prepared by the aforementioned methods, may be analyzed according to standard procedures.

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The present invention also provides antibodies capable of immunospecifically binding to MOP proteins of the invention. Polyclonal or monoclonal antibodies directed toward any of MOPs 1-9 may be prepared according to standard methods. Monoclonal antibodies may be prepared according to general methods of Köhler and Milstein, following standard protocols. In a preferred embodiment, antibodies have been prepared, which react immunospecifically with various epitopes of the MOPs.

Polyclonal or monoclonal antibodies that immunospecifically interact with MOPs can be utilized for identifying and purifying such proteins. For example, antibodies may be utilized for affinity separation of proteins with which they immunospecifically interact. Antibodies may also be used to immunoprecipitate proteins from a sample containing a mixture of proteins and other biological molecules. Other uses of anti-MOP antibodies are described below.

IV. Uses of MOP-Encoding Nucleic Acids, MOP Proteins and Anti-MOP Antibodies

A. MOP-Encoding Nucleic Acids

MOP-encoding nucleic acids may be used for a variety of purposes in accordance with the present invention. MOP-encoding DNA, RNA, or fragments thereof may be used as probes to detect the presence of and/or expression of genes encoding MOPs. Methods in which MOP-encoding nucleic acids may be utilized as probes for such assays include, but are not limited to: (1) in situ

hybridization; (2) Southern hybridization (3) northern hybridization; and (4) assorted amplification reactions such as polymerase chain reactions (PCR). In addition, recombinant cellular assay systems to examine signal transduction pathways in which the MOPs are involved are described below.

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The MOP-encoding nucleic acids of the invention may also be utilized as probes to identify related genes either from humans or from other species. As is well known in the art, hybridization stringencies may be adjusted to allow hybridization of nucleic acid probes with complementary sequences of varying degrees of homology. Thus, MOP-encoding nucleic acids may be used to advantage to identify and characterize other genes of varying degrees of relation to the respective MOPs, thereby enabling further characterization the AHR or related signaling cascades. Additionally, they may be used to identify genes encoding proteins that interact with MOPs (e.g., by the "interaction trap" technique, or modifications thereof, as described in Example 1), which should further accelerate elucidation of these cellular signaling mechanisms.

Nucleic acid molecules, or fragments thereof, encoding MOPs may also be utilized to control the production of the various MOPs, thereby regulating the amount of protein available to participate in cellular signaling pathways. In one embodiment, the nucleic acid molecules of the invention may be used to decrease expression of certain MOPs in cells. In this embodiment, full-length antisense molecules are employed which are targeted to respective MOP genes or RNAs, or antisense oligonucleotides, targeted to specific regions of MOP-encoding genes that are critical for gene expression, are used. The use of antisense molecules to decrease expression levels of a pre-determined gene is known in the art. In a preferred embodiment, antisense oligonucleotides are modified in various ways to increase

their stability and membrane permeability, so as to maximize their effective delivery to target cells in vitro and in vivo. Such modifications include the preparation of phosphorothicate or methylphosphonate derivatives, among many others, according to procedures known in the art.

In another embodiment, the transcription regulation activity of bHLH-PAS homodimers or heterodimers involving MOPs may be blocked by genetically engineering a target cell to express a defective MOP - specifically one that has been modified to be unable to bind DNA. When the defective MOP dimerizes, the dimer is also unable to bind DNA, and therefore is unable to carry out its transcriptional regulatory function.

In another embodiment, overexpression of various MOPs is induced, which can lead to overproduction of a selected MOP. Overproduction of MOPs may facilitate the isolation and characterization of other components involved in protein-protein complex formation occurring during the MOP-related signal transduction in cells.

As described above, MOP-encoding nucleic acids are also used to advantage to produce large quantities of substantially pure MOP proteins, or selected portions thereof.

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B. MOP Proteins and Anti-MOP Antibodies

Purified MOPs, or fragments thereof, may be used to produce polyclonal or monoclonal antibodies which also may serve as sensitive detection reagents for the presence and accumulation of MOPs (or complexes containing the MOPs) in cultured cells or tissues or in intact organisms. Recombinant techniques enable expression of fusion proteins containing part or all of a selected MOP protein. The full length protein or fragments of the protein may be used to advantage to generate an array of monoclonal or polyclonal antibodies specific for various epitopes of the protein, thereby providing even greater sensitivity for

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detection of the protein in cells or tissue.

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Polyclonal or monoclonal antibodies immunologically specific for a MOP may be used in a variety of assays designed to detect and quantitate the protein. Such assays include, but are not limited to: (1) flow cytometric analysis; (2) immunochemical localization of a MOP in cells or tissues; and (3) immunoblot analysis (e.g., dot blot, Western blot) of extracts from various cells and tissues. Additionally, as described above, anti-MOPs can be used for purification of MOPs (e.g., affinity column purification, immunoprecipitation).

C. Recombinant Cells and Assay Systems

Genetically engineered cells, such as yeast cells or mammalian cells, may be produced to express any 15 one, or a combination, of MOPs described herein. cells can be used to evaluate the binding interactions between MOPs, or between a MOP and another member of the bHLH-PAS superfamily (e.g., AHR, ARNT), and the 20 requirement for homodimerization or heterodimerization of the MOPs for initiation of transcriptional control of a reporter gene driven by appropriate enhancer elements. In addition, such recombinant cells can be used to study the effect of external stimuli, such as hypoxia or TCDD, on activation of a selected MOP, or they can be used to 25 screen panels of drugs for control of MOP-involved signal transduction pathways. U.S. Patent No. 5,650,283 to Bradfield et al., the disclosure of which is incorporated herein by reference, describes recombinant cellular systems and assays for detecting agonists to the AHR. 30 These materials and methods may be used similarly to design recombinant systems for evaluating any of MOP1-MOP8, in the presence or absence of an external stimulant.

Appropriate yeast cells for production of such recombinant systems include Saccharomyces cerevisiae and Saccharomyces pombe. Yeast strains carrying endogenous functional HSPs may be utilized (e.g., A303 obtained from

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Rick Gaber, Northwestern University, or commercially available equivalents). Yeast strains in which the genes encoding HSPs have been disrupted may also be utilized (e.g., GRS4, obtained from Susan Lindquist, University of Chicago), affording an opportunity to examine the relationship of various MOPs to HSPs.

Appropriate mammalian cells for production of such recombinant systems include COS, Hep3b, HepGr and Hepalclc7 cells, among others.

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In one type of assay where the MOP signal transduction pathway is affected by an external stimulus (i.e. an agonist such as TCDD in the AHR-ARNT system, or cobalt chloride in the MOP1/HIF1 α -ARNT system), an appropriate cell can be transformed with an expression plasmid expressing a full length agonist receptor MOP, along with its dimerization partner (if the MOP forms heterodimers) and a reporter plasmid expressing a reporter gene, such as LacZ or luciferin, which is driven by an appropriate enhancer element. The presence or potency of a selected agonist may be determined by its ability to activate transcription of the reporter gene in the recombinant system.

In another embodiment, a recombinant system that does not rely on heterodimerization can be constructed.

In this case, a cell is transformed with an expression plasmid expressing a chimeric agonist-sensitive MOP, along with a reporter plasmid expressing a reporter gene driven by a suitable promoter. The chimeric MOP is modified to replace the heterodimerization domains (i.e. the bHLH-PAS domain) with a DNA binding domain, such as LexA or Gal4. Such chimeras will homodimerize and activate transcription of genes positioned downstream of LexA or Gal4 binding sites engineered into the reporter plasmid.

In a preferred embodiment, described in detail
in Example 1, a modified yeast "two hybrid" system is used
to assess binding interactions between MOPs (and other
bHLH-PAS proteins) and the subsequent initiation of

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transcriptional control. For instance, as described in Example 1, fusion proteins were constructed in which the DNA binding domain of the bacterial repressor, LexA, was fused to the bHLH-PAS domains of the MOP proteins.

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in Example 3.

Interactions were tested by cotransformation of each LexAMOP construct with either the full length AHR or ARNT into the L40 yeast strain, which harbors an integrated lacZ reporter gene driven by multiple LexA operator sites. In this system, LexAMOP fusions which interact with AHR or ARNT drive expression of the lacZ reporter gene. The effect of various agonists on reporter gene expression can also be evaluated using this system.

Any one or more of the aforementioned recombinant cell systems and assays can be used to screen panels of drugs for their effect on specific signal transduction pathways. For instance, recombinant systems employing any or MOPs 1, 2, 6 or 7 may be used to screen for drugs that stimulate red blood cell synthesis, angiogenesis or glucose metabolism.

20 Recombinant systems employing any of MOPs 3, 4, 8 or 9 may be used to screen for drugs that modify circadian rhythms. In connection with this embodiment, as described in greater detail in Example 2, we have determined the binding sequence for the MOP3/MOP4 heterodimer, and have constructed the following 25 recombinant plasmids: PL833, a MOP3 expression vector for mammalian cells; PL834, a MOP4 expression vector for mammalian cells; and PL880, a reporter plasmid (expressing luciferase) driven by the MOP3/MOP4 consensus enhancer 30 sequence GCA CACGTG ACC. When the three plasmids are introduced into a mammalian cell, the reporter gene responds to the presence of the MOP3/MOP4 dimer. system is used in a high throughput microwell assay to screen for compounds that are specific activators or 35 inhibitors of these transcription factors. A similar system has been established for MOP7 (HIF3 α), as set forth

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The following examples are intended to illustrate embodiments of the invention. They are not intended to limit the scope of the invention in any way.

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EXAMPLE 1 Identification and Characterization of MOPs 1-5 cDNAs and Encoded Proteins

We employed an iterative search of human expressed sequence tags to identify novel basic-helix-loop-helix-PAS (bHLH-PAS) proteins that might interact with either the Ah receptor (AHR) or the Ah receptor nuclear translocator (ARNT). In this example, we describe the identification and characterization of five new "Members of the PAS superfamily," or MOPs 1-5, that are similar in size and structural organization to the AHR and ARNT.

20 METHODS

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Search Strategy. The bHLH-PAS domains of the huAHR, huARNT, drSIM, and the PAS domain of drPER were used as query sequences in BLASTN searches of the GenBank database between December of 1994 and October of 1995, using the following default values: Database = NR, expect = 10, word length = 12 (Altschul et al., J. Mol. Biol. 215: 403-410, 1990). Preliminary experiments comparing AHR and PER led us to define candidate ESTs as those "hits" that yielded scores of 150 or higher. As a method to confirm the similarity of these EST sequences to known bHLH-PAS proteins, each candidate EST was subsequently compared to the NR subset of GenBank using the BLASTX program, matrix = blosum 62, word length = 3. Only ESTs that retrieved known bHLH-PAS proteins by this method of confirmation were further characterized.

Oligonucleotide Sequences: Sequences of oligonucleotides are given below. In cases where the oligonucleotide was used in gel shift assays, the 6 bp target sequence is underlined.

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OL21 5' CGAGGTCGACGGTATCG 3'
     OL22 5' TCTAGAACTAGTGGATC 3'
     OL124 5' CCCAAGCTTACGCGTGGTCTTTGAAGTCAACCTCACC 3'
     OL145 5' AGCTCGAAATTAACCCTCACTAAAGG 3'
     OL176 5' CGGGATCCTTACACATTGGTGTTGGTACAGATGATGTACTC 3'
     OL180 5' GCGTCGACTGATGAGCAGCGGCGCCAACATCACC 3'
     OL201 5' GATAAGAATGCGGCCGCAGATCTGGGTCCGAAGCACACG 3'
     OL202 5' CATTACTTATCTAGAGCTCG 3'
     OL226 5' CGGGATCCTCATGGCGGCGACTACTGCCAACC 3'
     OL365 5' GACAGTTGCTTGAGTTTCAACC 3'
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     OL386 5' TTATGAGCTTGCTCATCAGTTGCC 3'
     OL387 5' CCTCACACGCAAATAGCTGATGG 3'
     OL392 5' CCGCTCGAGTGATGAGCAGCGGCGCCAACATCACC 3'
     OL393 5' CCGCTCGAGTGGCAGCTACAGGAATCCACC 3'
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     OL404 5' GCGGTACCGGGACCGATTCACCATGGAG 3'
     OL414 5' TCGAGCTGGGCAGGCTACGTGGCAAGGC 3'
     OL415 5' TCGAGCCTTGCCACGTACCCTGCCCAGC 3'
     OL418 5' GTAAAACGACGGCCAGT 3'
     OL419 5' GGAAACAGCTATGACCATG 3'
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     OL443 5' TCGAGCTGGGCAGGGTGCGTGGCAAGGC 3'
     OL444 5' TCGAGCCTTGCCACGCACCCTGCCCAGC 3'
     OL445 5' TCGAGCTGGGCAGGTCACGTGGCAAGGC 3'
     OL446 5' TCGAGCCTTGCCACGTGACCTGCCCAGC 3'
     OL447 5' TCGAGCTGGGCAGGTTGCGTGGCAAGGC 3'
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     OL448 5' TCGAGCCTTGCCACGCAACCTGCCCAGC 3'
     OL450 5' TACTGGCCACTTACTACCTGACC 3'
     OL456 5' AACCAGAGCCATTTTTGAGACT 3'
     OL477 5' GCTCTAGAGGCCACAGCGACAATGACAGC 3'
     OL479 5' GATCGGAGGTGTTCTATGAGC 3'
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     OL489 5' TTAGGATGCAGGTAGTCAAACA 3'
     OL496 5' GTTCTCCATGGACCAGACTGA 3'
     OL499 5' CGGGTACCCTGGGCCCTACGTGCTGTCTC 3'
     OL500 5' CGGCTAGCCTCTGGCCTCCCTTGATGA 3'
     OL514 5' CTGGGAGCCTGCCTTCA 3'
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    OL520 5' CCCAAGGAGAGGCGTGAT 3'
     OL540 5' GGGATCCTCGTCGCCACTG 3'
     OL541 5' ATGCAGTACCCAGACGGATTTC 3'
    OL560 5' TGCACGGTCACCAACAGAG 3'
    OL561 5' TTGCCAGTCGCATGATGGA 3'
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     OL565 5' CTGAACAGCCATCCTTAG 3'
    OL568 5' AGCTTGCCCTACGTGCTGTCTCAG 3'
    OL569 5' AATTCTGAGACAGCACGTAGGGCA 3'
    OL590 5' AGAGGTGCTGCCCAGGTAGAA 3'
    OL611 5' CAATGATGAGGGAAACACTG 3'
    OL657 5' CGGGATCCCGTCAACTGGAGATGAGCAAGGAG 3'
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     OL665 5' CTGCAAAAATCCGATGACCTCTT 3'
     OL681 5' CGGGCAGCAGCGTCTTC 3'
    OL682 5' GCGTCCGCAGCCCCAGTTG 3'
    OL683 5' TTCAATGTTCTCATCAAAGAGC 3'
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     OL684 5' GAACAGTTTTATAGATGAATTGGC 3'
     OL689 5' GAGGTGTTTCAATTCATCGTCT 3'
    OL715 5' GGGATCCGTGACCGATTCACCATGGAG 3'
    OL716 5' CTGCAGGTCACACAACGTAATTCACACA 3'
    OL717 5' GGGATCCGTATGACAGCTGACAAGGAG 3'
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     OL718 5' GGTCGACGTCACAGGACGTAGTTGACACA 3'
     OL719 5' GAATCCATGAGCAAGGAGGCCGTG 3'
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OL720 5' GGTCGACGTCAAACAACAGTGTTAGTTGA 3' OL721 5' GGGATGCGTATGGATGAGATGAGAAAGAC 3' OL722 5' GGTCGACGCTAGACCGAGTGTGTGCA 3'

5 Cloning strategy. To obtain extended open reading frames for each EST, an anchored-PCR strategy was employed to amplify additional flanking sequence from a variety of commercial cDNA libraries that were constructed in the phagemid Lambda Zap (Tissues; HepG2, Fetal Brain 10 and Skeletal Muscle; Stratagene, La Jolla, CA) (Table 1) (Innis et al. (eds), PCR Protocols: a Guide to Methods and Applications, Academic Press, San Francisco, 1990). resulting PCR products were subjected to agarose gel electrophoresis, transferred to a nylon membrane and 15 analyzed by hybridization with a 32P-labeled probe generated from the corresponding parent EST plasmid (Table After autoradiography, the positive PCR products were purified by gel electrophoresis and cloned using the pGEM-T vector system (Promega, Madison, WI). 20 sequencing was performed to characterize each positive clone.

TABLE 1

MOP cDNA Clone Information

Row 1: clones (in parentheses) containing the candidate ESTs were requested from their laboratory of origin.

Row 2: the Genbank accession number for each original EST is indicated.

Row 3: oligonucleotides used in library screening. Sequence information generated from this original clone was used to design oligonucleotides for use in an anchored PCR strategy, whereby gene-specific and vector-specific primers were used to amplify 5' and 3' portions of the cDNA. Vector specific primers corresponded to modified T3 (5', OL145) or T7 (3', OL146) primers. A matrix of genespecific primers against annealing temperature (50-65°C) was attempted for each clone, generally leading to at least one successful reaction.

Row 4: the cDNA libraries from which additional sequence of positive clones was identified.

Row 5: size of ORFs. We define a complete ORF by the presence of an in-frame stop codon 5' to a methionine codon that lies within a Kozak consensus sequence for translational initiation. The 3' end of open reading frames are defined by the presence of an inframe termination codon. An asterisk (*) denotes a clone which does not meet these criteria (see text).

Row 6: Genbank accession numbers of the final MOP cDNAs

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are given.

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	MOP1/HIF1α	MOP2	мор3	MOP4	MOP5
Laboratory of origin (clone desig.)	Bell (hbc025)	IMAGE (67043)	IMAGE (23820, 50519)	Liew (F9047, PL420)	IMAGE (42596)
EST Genbank accession number	T10821	T70415	Т77200, Н17840	R58054	R67292
Gene- specific oligo used in PCR	OL365 (5')	OL456 (5') OL496 (3') OL514 (3') OL541 (3')	OL489 (5')	OL520 (5')	OL540 (5')
Library screened	НерG2	HepG2	Fetal brain	HeLa	HepG2
ORF size	826	870	624	642*	412*
Final cDNA Genbank accession number	U29165	U51626	U51627	U51625	U51628

Plasmid Construction for Expression In Vivo.

Sequence information from each EST was used to design PCR primers for the amplification of cDNA from commercially available libraries. Expression plasmids were constructed by standard protocols (Sambrook et al., 1989). For a summary of clone designations, PCR primers, DNA templates and GenBank accession numbers, refer to Table 1. A brief description follows.

MOP1 expression vectors. Oligonucleotides OL404 and OL365 were used as primers in a PCR to amplify a 970 bp fragment from a HepG2 cell cDNA library. This fragment was cloned into the pGEM-T vector in the T7 orientation and designated PL439. To generate pGMOP1, the Sall/XhoI

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fragment of hbc025 was subcloned into SalI digested PL439. To increase transcription efficiency of the MOP1 cDNA, pGMOP1 was digested with KpnI and SacI and this fragment subcloned into the corresponding sites of pSputk generating PL415 (Stratagene, La Jolla, CA) (Falcone & Andrews, Mol. Cell. Biol. 11: 2656-2664, 1991). The complete ORF of the MOP1 cDNA was amplified using the PCR and oligonucleotides OL425 and OL536. This fragment was digested with BamHI and ligated into the BamHI site in the pSport polylinker (Life Technologies, Inc.). This plasmid was designated PL611.

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MOP2 expression vectors. PCR was employed using OL477 and OL450 to amplify a 931 bp MOP2 fragment from a HepG2 cDNA library. This fragment was cloned into pGEM-T in the SP6 orientation and designated PL424. Using OL560 and OL590, PCR amplification from this same library yielded a 3' fragment of the MOP2 cDNA. This fragment was cloned into pGEM-T in the SP6 orientation and was designated PL445. PL424 was digested SalI and EcoRI and the fragment ligated into a SalI/EcoRI digested PL445 to generate a full ORF MOP2 expression vector designated PL447. The complete ORF of the MOP2 cDNA was cloned into pSport as follows; PL447 was digested with SacII, treated with the Klenow fragment of DNA Polymerase I in the presence of dNTPs, and subsequently digested with Sall. This fragment was purified and ligated into pSport digested with HindIII, repaired with Klenow, then digested with Sall. This construct was designated PL477.

MOP3 expression vectors. Using the primers

OL145 and OL489 and a human fetal brain cDNA library as template, the PCR was used to obtain a 1380 bp fragment. This fragment was isolated and cloned into pGEM-T as above, and this plasmid designated PL487. A fragment of MOP3 was obtained by the PCR using Pfu polymerase

(Stratagene), primers OL657 and OL689 and PL487 as template. To obtain a full length MOP3 cDNA fragment, the megaprimer fragment obtained above was used in the PCR

against oligonucleotide OL611 using IMAGE clone 50519 as a template (Sarkar & Somers, Biotechniques 8: 404-407, 1990). This product was cloned into pGEM-T in the SP6 orientation as above and designated PL425.

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MOP4 expression vectors. Using primers OL520 and OL145 and a HepG2 cDNA library as template, the PCR was performed to isolate a 5' fragment of the MOP4 cDNA. This fragment was cloned in the T7 orientation of pGEM-T and designated PL448. The cDNA insert of the phage clone F9047 (C.C. Liew, Toronto, CA) was amplified by the PCR using oligonucleotides OL418 and OL419 and subcloned into the pGEM-T vector (Hwang et al., J. Mol. Cell. Cardiol. 26: 1329-1333, 1994). This clone was designated PL420. An EcoRI fragment of PL448 was isolated and cloned into a partially EcoRI digested PL420. This clone was subjected to the PCR with oligonucleotides OL698 and OL146, and the fragment cloned into the pGEM-T vector, and designated PL545.

MOP5 expression vectors. The PCR was used to obtain a 1260-bp fragment of the MOP5 gene using oligonucleotides OL685 and OL686 and IMAGE clone 42596 as template. This fragment was purified and subcloned into the pGEM-T vector as above in the SP6 orientation. This plasmid was designated PL528 and subsequently digested with SalI and partially digested with NcoI. This fragment was ligated into Nco I/SalI-cut pSputk, and the resulting vector designated PL554.

Rypoxia responsive luciferase reporters. The plasmid pGL2EPOEN was constructed as follows: The hypoxia responsive enhancer from the 3' region of the EPO gene was amplified by PCR using oligonucleotides OL499 and OL500 and human genomic DNA as template (amplified fragment corresponds to nucleotides 127 to 321 as reported in the EPO structural gene sequence found in GenBank Accession GBL16588). This fragment was digested with KpnI and NheI and cloned into the corresponding sites of the plasmid pGL2-Promoter (Promega).

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Antibody Production. Antisera against MOP1, MOP2, AHR and ARNT were prepared in rabbits using immunization protocols that have been described previously (Poland et al., Mol. Pharmacol. 39: 20-26, 1991; Pollenz et al., Mol. Pharmacol. 45: 428-438, 1994). Crude 5 antisera was chosen for use in all coimmunoprecipitation experiments and the PI sera from the same rabbit served to preclear the samples. For MOP1, the plasmid hbc025 was digested with EcoRI and the 604 bp fragment was treated 10 with the Klenow fragment of DNA polymerase-1 in the presence of dNTPs and cloned into the SmaI site of the histidine tag fusion vector pQE-32 (Qiagen, Chatsworth, CA). This clone, designated PL377, was transformed by electroporation into M15(REP4) cells for expression under IPTG induction. The expressed protein was purified from 8 15 M urea using Ni-NTA agarose, extensively dialyzed against 25 mM MOPS, pH 7.4, 100 mM KCl, and 10% glycerol before its use as an immunogen. For AHR, the human cDNA clone PL71 (Dolwick et al., Mol. Pharmacol. 44: 911-917, 1993) was digested with BamHI and cloned into the corresponding 20 site of the histidine fusion vector pQE31 (Qiagen). AHR protein fragment was expressed and purified exactly as described for MOP1 (above). Antiserum produced against this protein was designated R2891. For MOP2 a SacI/PstI fragment of PL445 was cloned into SacI/PstI cut pQE-31 to 25 generate PL456. This clone, designated PL456, was transformed into M15(REP4) cells and the protein expressed under IPTG induction. The histidine tagged fusion protein was first extracted in guanidine hydrochloride, dialyzed 30 extensively and purified on Ni-NTA agarose as above. Antiserum produced against this protein was designated R4064. ARNT-specific antisera was raised against huARNT protein purified from baculovirus as previously described (Chan et al., J. Biol. Chem. 269: 26464-26471, 1994).

North rn Protoc 1. Multiple tissue northern blots containing 2 μ g of poly(A)+ mRNA prepared from human heart, brain, placenta, lung, liver, skeletal muscle,

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kidney, and pancreas were probed with random primed cDNA fragments using an aqueous hybridization protocol (Clontech, Palo Alto, CA). Hybridization solution contained 5 X SSPE (0.75 M NaCl, 50 mM NaH, PO, 5 mM Na₂EDTA, pH 7.4) 2 X Denhardt's solution (0.04 % w/v Ficoll 400, 0.04% w/v polyvinylpyrrolidone, 0.04 % w/v Bovine Serum Albumin), 0.5 % SDS, and 100 ug/mL heat denatured salmon sperm DNA. A blot was prehybridized for 3-6 hours at 65°C, the hybridization solution was changed 10 and 1-5x10⁶ cpm/mL of a random primed cDNA fragment was Samples were hybridized overnight at 65°C, washed twice with 2 X SSC (0.3 M NaCl, 30 mM Na₃Citrate, pH 7.0), 0.5 % SDS at room temperature, once with 1% SSC, 0.1% SDS at the hybridization temperature, and once with 0.1 X SSC, 15 0.1% SDS at the hybridization temperature.

Yeast Two-Hybrid Analysis. A modified yeast interaction trap was employed to identify those MOPs that could interact with the AHR or ARNT. LexAMOP chimeras were constructed to fuse the bHLH-PAS domains of the MOP proteins with the DNA binding domain of the bacterial 20 protein LexA (amino acids 1-202) (Bartel et al., BioTechniques 14: 920-924, 1993). To amplify the region corresponding to the bHLH-PAS domains of MOP1, OL715 and OL716 were employed in the PCR using PL415 as template. 25 To amplify the region corresponding to the bHLH-PAS domains of MOP2, OL717 and OL718 were employed in the PCR using PL447 as template. To amplify the region corresponding to the bHLH-PAS domains of MOP3, OL719 and OL720 were employed in the PCR using PL486 as template. To amplify the region corresponding to the bHLH-PAS 30 domains of MOP4, OL721 and OL722 were employed in the PCR using PL545. Since a more detailed domain map existed for the AHR, a construct was made with a fine deletion of the transactivation domain. The N-terminal portion of the AHR 35 was amplified by the PCR using oligonucleotides OL180 and OL124 and pmuAHR as template (Dolwick et al., Proc. Natl.

Acad. Sci. USA 90: 8566-8570, 1993). This product was

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digested with KpnI and SalI, and cloned into the corresponding sites of pSG424 (Sadowski & Ptashne, Nucl. Acids Res. 17: 7539, 1989). This clone was designated PL187. The 3' end of the AHR cDNA was amplified by PCR 5 using oligonucleotides OL201 and OL202 and pmuAHR as template. This product was digested with NotI, and cloned into the corresponding site of pSGAhN-delta-166 (Dolwick et al, Proc. Natl. Acad. Sci. USA 90: 8566-8570, 1993). This clone was designated PL188. PL188 was digested with 10 KpnI and XbaI, and this fragment cloned into the corresponding sites of PL187. This clone was designated A cDNA fragment of the AHR was amplified by the PCR using OL392 and OL393 and PL204 as template. product was cloned using the pGEM-T system, and designated pGTAHR-delta-TAD. This construct was digested with XhoI 15 and this fragment ligated into SalI cut pBTM116 (Vojtek et al., Cell 74: 205-214, 1993). This construct was designated pBTMAHR. LexAARNT was constructed by PCR using oligonucleotides OL226 and OL176 and PL87 as template. The PCR product was cloned into pGEM-T as above, and the 20 BamHI fragment cloned into a BamHI digested pGBT9 vector (Clontech). This construct was cut with BamHI and subcloned into a BamHI digested pBTM116. This construct was designated LexAARNT. Following amplification these 25 products were purified and cloned into the pGEM-T vector. These clones were designated PL537 (MOP1), PL538 (MOP2), PL539 (MO3), and PL540 (MOP4). These plasmids were digested with BamHI/PstI (PL537), BamHI/SalI (PL538 and PL540), and EcoRI/SalI (PL539), and these fragments 30 ligated into the appropriately digested pBTM116. clones were designated LexAMOP1, LexAMOP2, LexAMOP3, and LexAMOP4, respectively. Full length expression plasmids harboring the AHR and ARNT were constructed as follows: PL104 (pSporthuAHR) was cut with Smal, the insert purified 35 and subcloned into Smal site of pCW10, and this plasmid was designated PL317. This clone was digested with Smal and subcloned into a SmaI cut pRS305, and this clone

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designated pRSAHR. The ARNT cDNA (PL101) was digested with Not I and XhoI, and cloned into the corresponding sites of pSGBMX1. This plasmid was designated PL371, and subsequently was digested with NotI and XhoI, this 5 fragment cloned into the corresponding sites of pSGBCU11. This clone was designated PL574. The LexAMOP fusion protein constructs were cotransformed with a yeast expression vector containing the full length AHR or ARNT into L40, a yeast strain containing integrated lacZ and 10 HIS3 reporter genes. As controls, LexAAHR and LexAARNT were cotransformed with AHR and ARNT. The strength of interaction was visually characterized by X-Gal (5-bromo-4-chloro-3-indolyl-(-D-galactoside) plate assays, performed after three days growth on selective media 15 (Bohen et al, Proc. Natl. Acad. Sci. USA 90: 11424-11428, To provide quantitation of the interaction strength, multiple colonies from yeast harboring each bHLH-PAS combination colonies were grown overnight in liquid media. Liquid cultures were grown for 5 hours, and 20 assayed for lacZ activity using the Galacto-Light chemiluminescence reporter system (Tropix, Bedford, MA). To determine the effect of AHR agonists on these interactions, yeast were also grown on plates or in liquid culture with and without 1 μM βNF (Carver et al., J. Biol. 25 Chem. 269: 30109-30112, 1994).

To ensure expression of each bHLH-PAS construct, western blot analysis was performed using antibodies raised against the LexA DNA binding domain. Yeast extracts were prepared from 15 mL overnight cultures derived from multiple colonies of yeast expressing each LexAMOP fusion protein. Cultures were subjected to centrifugation at 1200 x g for 5 minutes and the pellet was resuspended in 500 μ L of 6 M Guanidinium-HCl, 0.1 M Na-Phosphate Buffer, 0.01 M Tris pH 8.0. This suspension was transferred to a fresh eppendorf tube containing 500 μ L of acid washed glass beads (Sigma), and mixed on the max setting in a Bead-Beater (BioSpec, Bartlesville, OK)

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for 3 minutes at 4°C. The samples were cleared by centrifugation at 14,000 x g, and 400 μ L of supernatant was precipitated with 400 μ L of 10 % TCA on ice. After clearing by centrifugation at 14,000 x g for 20 minutes at 4°C, the extracts were resuspended in SDS loading buffer and subjected to SDS-PAGE analysis. Following electrophoresis, proteins were transferred to nitrocellulose membrane and detected with LexA antisera and secondary antibodies linked to alkaline phosphatase by standard protocols (Jain et al., J. Biol. Chem 269: 31518-31524, 1994).

pGL2EPOEN was cotransfected with pSport, PL464 (pSportMOP1), or PL477 (pSportMOP2) using the Lipofectin protocol (Life Technologies, Inc.). Briefly, the expression vector was mixed with the epo-reporter and the beta-galactosidase control plasmid pCH110 (Clontech) at a 3:1 charge ratio of TFX-50 reagent (Promega) and incubated for 15 minutes at room temperature. The lipofection media (200 μL) was added to Hep3b cells in 4 cm² plates in the presence of serum. The cells were incubated at 37°C for 2 hr. Following incubation, fresh media was added, and the cells were incubated for an additional 48 hours prior to harvesting. Cell extraction and beta-galactosidase assays were performed using the Galacto-Light assay according to manufacturer's protocols (Tropix).

Coimmunoprecipitation with Hsp90. Each MOP construct was in vitro translated in the presence of [35S]-methionine in a TNT coupled transcription/translation system (Promega). Hsp90 immunoprecipitation assays were performed with monoclonal antibody 3G3p90 or a control IgM antibody, TEPC 183 (Sigma) essentially as described. Each immunoprecipitation was subjected to SDS-PAGE, and the resulting gel was dried. The level of radioactivity in each coprecipitated protein band was quantified on a Bio-Rad GS-363 Molecular Imager System. The amount of protein immunoprecipitated with the Hsp90 antibody is

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presented as a percentage of the amount of murine AHR immunoprecipitated in parallel assays.

RESULTS

EST Search. Our initial BLAST searches in 5 December 1994 were performed with the bHLH-PAS or PAS domains of all family members known at that time (AHR, ARNT, SIM and PER). In these searches we identified an EST clone, hbc025, derived from human pancreatic islets (Table 1) (Takeda et al., Hum. Mol. Genet. 2: 1793-1798, 10 To confirm this similarity, we performed a BLASTX search, comparing hbc025 to the GenBank database and found that this sequence was most homologous to Drosophila SIM. This EST clone was designated MOP1. The bHLH-PAS domains 15 of all family members, including MOP1, were again searched from May to October of 1995. Human ESTs that recorded BLASTN scores above 150 were again retrieved and confirmed using the BLASTX algorithm. This routine resulted in the discovery of six ESTs with significant homology to the 20 bHLH-PAS domains of known members (Table 1).

cDNA Cloning. In order to more completely characterize the similarities and domain structures of the candidate clones, an anchored-PCR strategy was employed to obtain additional flanking cDNA sequence using phagemid libraries as a template. Comparison of amino acid 25 sequences of these bHLH-PAS proteins is displayed in Figure 2. Upon characterization of the open reading frames, it was learned that two of these ESTs (F06906 and T77200) corresponded to the same gene product (Table 1). 30 Thus, we designated these remaining five unique cDNAs as "Members of PAS superfamily" or MOPs 1-5. strategy provided what appeared to be the complete ORFs of MOP1, MOP2 and MOP3 based upon the following criteria: (1) at their 5' ends these clones contain an initiation 35 methionine codon (AUG) downstream of an in-frame stop codon, and (2) at their 3' ends these clones contain an in-frame stop codon followed by no obvious open reading

frames. In addition, the nucleotide sequences flanking of the MOP1 and MOP2 most 5' AUG codons (see GenBank accessions U29165 and U51626) are in reasonable agreement with the proposed optimal context for translational initiation, i.e., CCACCAUGG (Kozak, Cell 44: 283-292, 1986; Kozak, Nucl. Acids Res. 15: 8125-8132, 1987).

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Using the same anchored-PCR technique, we were unable to obtain the complete open reading frames of MOP4 or MOP5. This may have been due to the low copy number of these mRNAs in the tissues from which our PCR source cDNA was obtained (see below). We did identify a potential start methionine for MOP4 and the 3' stop codon for MOP5 (Figure 2). Our preliminary designation of the MOP4 start methionine is tentative and is based only on its proximity to the start methionines of MOP1, MOP2, MOP3, AHR and SIM (Figure 2) (Burbach et al., Proc. Natl. Acad. Sci. USA 89: 8185-8189, 1992; Nambu et al., Cell 67: 1157-1167, 1991). The fact that only one of the six nucleotides flanking the MOP4 AUG codon (ATTTAATGG) matches the consensus sequences for optimal translational initiation provides an indication that a more 5' initiation codon may exist. Therefore, the initiation codon of MOP4 is uncertain and that of MOP5 remains to be identified. This low level expression is consistent with our difficulties in amplifying these cDNAs by PCR (see above) and suggests that expression may be limited to specific cell types or developmental time periods not identified in our study.

Tissue Specific Expression. To characterize the tissue specific expression patterns of the MOP mRNAs,

Northern blots of poly A(+) RNA from eight human tissues were probed with random primed cDNA restriction fragments. Single transcripts of 3.6 kb (MOP1/HIF1α), 6.6 kb (MOP2) and 3.2 kb (MOP3) were detected. Expression levels of each mRNA varied significantly between tissues, with MOP1 being highest in kidney and heart, MOP2 highly expressed in placenta, lung, and heart, and MOP3 highly expressed in skeletal muscle and brain. No detectable message was

detected for MOP4 or MOP5 by our northern blot protocol.

Identification of Novel AHR or ARNT Partners:

1. Interaction of MOPs with the AHR and ARNT in vitro; Coimmunoprecipitation experiments. We first performed coimmunoprecipitation experiments to determine if MOPs 1-4 had the capacity to interact with either the AHR or ARNT in vitro. These proteins were expressed in a reticulocyte lysate system in the presence of ³⁵S-methionine and then incubated in the presence or absence of the AHR or ARNT.

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- Complex formation was assayed by coimmunoprecipitation with AHR or ARNT specific antisera, followed by quantitation of coimmunoprecipitated ³⁵S-labeled MOP by phosphoimage analysis. Interactions were identified by a reproducible increase in an AHR or ARNT-dependent
- precipitation of MOP protein. Because we have observed considerable variability in this coimmunoprecipitation assay, each experiment was performed at least three times.

In the AHR interaction studies, we observed that MOP3 was coimmunoprecipitated with AHR. The positive control, ARNT-AHR interaction, was also reproducible, but weaker. Neither MOP1, MOP2 or MOP4 could be shown to interact with the AHR by this protocol. The ARNT protein displayed a broad range of interactions and was shown to coimmunoprecipitate with AHR (positive control), MOP1 and MOP2, but not MOP3 or MOP4.

Interaction of MOPs with the AHR and ARNT in vivo;
Yeast two-hybrid experiments. To determine if MOP-AHR or
MOP-ARNT complexes could form in vivo, a modified
interaction trap was employed (Fields & Song, Nature 340:
245-246, 1989; Chien et al., Proc. Natl. Acad. Sci. USA
88: 9578-9582, 1991) (Figure 3). Fusion proteins were
constructed in which the DNA binding domain of the
bacterial repressor, LexA, was fused to the bHLH-PAS
domains of the MOP proteins (Fig. 3A). The bHLH-PAS
domains were chosen because they harbor both the primary
and secondary dimerization surfaces of this family of

proteins and they do not harbor transcriptional activity

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that would interfere with this assay (Jain et al., 1994, supra). Interactions were tested by cotransformation of each LexAMOP construct with either the full length AHR or ARNT into the L40 yeast strain, which harbors an integrated lacz reporter gene driven by multiple LexA operator sites. In this system, LexAMOP fusions which interact with AHR or ARNT drive expression of the lacz reporter gene.

We assessed the relative strength of these interactions by both a direct lacZ plate assay and by 10 quantitation of the reporter activity in a liquid culture (Figure 3). In all cases, these two methods of detection were equivalent. To test the validity of this model system as a method to detect bHLH-PAS interactions. 15 LexAAHR and LexAARNT constructs were cotransformed with either the full length ARNT or AHR. In these control experiments, we were able to demonstrate the specificity of AHR-ARNT interaction and its dependence on the presence of the agonist βNF . The LexAAHR-ARNT interaction in the 20 presence of BNF was 913 fold above background, while the LexAARNT-AHR interaction in the presence of BNF was 14 fold above background. Both combinations showed ligand inducibility. The LexAAHR-ARNT interaction in the presence of BNF was 6.4 fold over LexAAHR-ARNT in the 25 absence of ligand, while the LexAARNT-AHR interaction in the presence of BNF was 2.0 fold over LexAARNT-AHR in absence of ligand. Despite our ability to readily detect the agonist-induced LexAARNT-AHR interaction in the two hybrid system, we were unable to detect any LexAMOP that 30 could interact with the AHR. That is, none of the LexAMOP fusion proteins appeared to interact with cotransformed AHR and drive lacz expression in the absence or presence of ligand.

Two of four MOP proteins tested were found to
interact with ARNT in the two hybrid assay. Both the
LexAMOP1 and LexAMOP2 interactions with full length ARNT
were extremely robust, 36 and 28 fold above background,

respectively (Fig. 3B). When compared with the LexAAHR-ARNT interaction in the presence of BNF, the LexAMOP1-ARNT and LexAMOP2-ARNT interactions were 24% and 69% as intense. These differences in LexAMOP1-ARNT and 5 LexAMOP2-ARNT interaction could be attributed to differences in expression levels or to subtle differences in vector construction. To control for relative expression of the LexAMOP fusions, protein extracts were prepared and western blot analysis was performed with LexA 10 specific antisera. We observed expression for each LexAMOP fusion proteins, indicating that negative results with LexAMOP3 and LexAMOP4 are not due to lack of expression.

3. DNA binding and specificity. Prompted by the observation that MOP1 and ARNT and MOP2 and ARNT 15 specifically interact, we next examined the ability of MOP-ARNT dimeric complexes to bind those DNA response elements recognized by other bHLH-PAS protein complexes. Reports from a number of laboratories have demonstrated 20 that bHLH-PAS dimers can bind to a variety of DNA elements: "DRE," TNGCGTG (Denison et al., J. Biol. Chem. 264: 16478-16482, 1989); "CME," ACGTG (Wharton et al., Development 120: 3563-3569, 1994); "SAE," GTGCGTG (Swanson et al., J. Biol. Chem. 270: 26292-26302, 1995); and "E-25 box," CANNTG (Sogawa et al., Proc. Natl. Acad. Sci. USA 92: 1936-1940, 1995; Swanson et al., 1995, supra). a gel-shift assay, we observed that MOP1-ARNT complexes specifically bound CACGTG and TACGTG, while the complex failed to bind GTGCGTG, TTGCGTG, and a non-palindromic Ebox, CATGTG. Previous reports have demonstrated that ARNT 30 homodimers are capable of binding the CACGTG sequence in vitro, and that this complex can drive reporter gene expression in vivo (Sogawa et al., 1995 supra, Swanson et al., 1995, supra). Our results suggest that the MOP1-ARNT dimeric complex binds the CACGTG oligonucleotide with a 35 higher affinity than either MOP1 or ARNT alone. MOP1 failed to form a productive DNA binding complex with the

AHR with any of the bHLH-PAS family response elements. As a comparison of MOP1-ARNT and MOP2-ARNT DNA binding, we provide results from gel shift assays using a double-stranded oligonucleotide containing a core TACGTG hexad binding site. WE observed that both MOP1-ARNT and MOP2-ARNT bound the TACGTG-containing oligonucleotide with approximately equal capacity and neither ARNT, nor MOP1, nor MOP2 could bind this DNA sequence alone. As additional controls, we confirmed the presence of the MOP proteins in the complex by showing that antisera raised against these proteins retarded the mobility of the complex.

- 4. Interaction of MOPs with Hsp90. In an effort to assess a MOP's ability to interact with Hsp90, we performed coimmunoprecipitation assays with anti Hsp90 antibodies. Given the remarkable stability of the Hsp90 complex with the AHR from the C57BL/6J mouse, we used this receptor species as a reference and compared all interactions relative to it. As additional controls, we
- immunoprecipitated ARNT and the human AHR as negative and positive controls, respectively. Despite our ability to readily detect huAHR-Hsp90 interactions, we were unable to detect ARNT, MOP2 or MOP5 interactions with Hsp90. In contrast, huAHR, MOP1, MOP3 and MOP4 all
- immunoprecipitated with HSP90-specific antisera. MOP3 formed the tightest interaction with HSP90, followed by the huAHR, MOP4 and MOP1 (71%, 53%, 31% and 17%, respectively).

30 DISCUSSION

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Since cDNAs encoding the complete open reading frames for MOPs 1-3 were available, most of the studies described in this example focused on those proteins. MOP4 was also included in some studies since our clone contained the sequences involved in dimerization, transcriptional activation and DNA binding of other bHLH-PAS proteins. Given the limited sequence information

on MOP5, this clone was typically not included in functional studies.

Tissue specific expression. We observed that each MOP mRNA displayed a unique tissue specific 5 distribution with MOP1 being highest in kidney and heart, MOP2 highly expressed in placenta, lung, and heart, and MOP3 highly expressed in skeletal muscle and brain. Previous studies conducted in our laboratory indicated that ARNT is expressed highly in skeletal muscle and 10 placenta, while the AHR is most prevalent in placenta, lung, and heart (Carver et al., 1994, supra; Dolwick et al., 1993, supra). The observation that these bHLH-PAS proteins are coexpressed in a variety of tissues supports the idea that cross talk between these signaling pathways 15 may be occurring in vivo and that multiple tissue specific interactions may be taking place. We also observed that AHR and MOP2 have very similar expression profiles in human tissues. An additional and equally important interpretation of these unique MOP expression profiles is 20 that unidentified partners exist for these bHLH-PAS proteins and that they regulate a number of undescribed biological pathways.

Interactions. Our interaction screening strategy was based on the large amount of functional data and the detailed domain maps available for the AHR and ARNT. An important assumption used in the design and interpretation of our studies is that some of the MOPs may be constitutive interactors in vivo (like ARNT) and others may be conditional interactors that require activation in order to dimerize in vivo (like AHR). We chose to employ coimmunoprecipitation as an initial interaction screen for a number of reasons. First, AHR and ARNT-specific antibodies are available that have been shown to pull down AHR-ARNT complexes. This suggests that if MOP-AHR or MOP-ARNT interactions occurred in vitro, that these same antibodies would recognize and pull down such complexes. Second, data from a number of laboratories, using

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independently derived antibodies indicates that coimmunoprecipitation of AHR-ARNT complexes is independent of AHR-ligand. This observation suggests that AHR or ARNT interactions with conditional MOP proteins might still be identified by coimmunoprecipitation even in the absence knowledge about how to activate a conditional MOP (e.g., identification of a ligand).

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As a secondary screen to characterize interacting MOPs, we employed a yeast interaction trap commonly referred to as the "Two Hybrid Assay". for use of this system comes from our previous observation that LexA-AHR chimeras are functional in yeast and provide a good model of AHR signaling and ARNT interaction. addition, this method provides an independent confirmation of those interactions identified by coimmunoprecipitation and also provides a demonstration that interactions can occur in vivo. One major limitation of this system is that it may be insensitive at detecting conditional MOPs that require activation prior to dimerization. An example of this can be seen with the AHR and ARNT. In the absence of ligand, the AHR appears to reside primarily in the cytosol and ARNT appears to be primarily nuclear. compartmentalization appears to be part of a cellular mechanism to prevent interaction of these proteins and minimize constitutive activity of the complex. important to point out that compartmentalization is only one component of AHR regulation, since ligand dependent DNA binding does occur in vitro in the presence of ARNT. Nevertheless, in vivo systems like the Two Hybrid Assay may yield false negative results for conditional MOP protein interactions that require an upstream activation event prior to nuclear transloction.

In light of the above considerations, our interpretation of the coimmunoprecipitation and two hybrid interaction results were as follows: First, since the MOP1-ARNT and the MOP2-ARNT interactions were confirmed in two independent systems these interactions should be

pursued further (see below). Second, the observation that MOP3 interact with the AHR in vitro, but not in vivo. suggests that MOP3 may be a conditional MOP that has the capacity to interact with the AHR in vivo. This idea gained support from Hsp90 interaction studies (below). Third, the suspicion that MOP3 is a conditional bHLH-PAS protein, coupled with the observation that MOP3 and AHR have the disparate expression profiles led us to delay study of this interaction until we learn how to activate MOP3 or until we have evidence that these two proteins are expressed in the same cell type. Fourth, our observation that ARNT can form dimers with two out of four MOPs examined suggests that ARNT is a highly promiscuous bHLH-PAS partner that may be a focus of cross talk between different MOP signaling pathways. The multiplicity of ARNT partnerships is supported by recent observations from a number of laboratories (Sogawa et al, 1995, supra; Swanson et al., 1995, supra).

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MOP1 and MOP2 interactions with ARNT. concordance of the coimmunoprecipitation and two hybrid 20 data led us to pursue the MOP1-ARNT and MOP2-ARNT interactions further. Given the pairing rules deduced from the interaction studies described above, we next attempted to determine if the MOP1-ARNT and MOP2-ARNT 25 complexes bound specific DNA sequences in vitro. reports indicated that the basic region of each bHLH partner generates specificity for a distinct DNA half-site of at least 3 bp. Data from a number of laboratories has indicated that the ARNT protein displays specificity for 30 the 3' GTG half site of the hexad target sequence, 5'NNCGTG3', where 5'NNC is the half site of the ARNT partner. To determine the half site specificity of the MOP1 protein when complexed with ARNT, we used gel shift analysis with oligonucleotides containing substitutions at the two variable 5' positions of 5'NNCGTG3'. 35 preliminary experiments indicated that MOP1-ARNT complex had greatest affinity for the 5'CAC and 5'TAC half sites.

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Because the MOP1 and MOP2 basic regions differed by only one amino acid residue and since this residue is not thought to be in a DNA contact position, we hypothesized that MOP2 would bind the same DNA sequences. To confirm this, we performed MOP2-ARNT gel shift assays using a double stranded oligonucleotide containing a core TACGTG hexad binding site. We observed that both MOP1-ARNT and MOP2-ARNT bound the TACGTG containing oligonucleotide, that neither MOP1 nor MOP2 could bind this sequence in the absence of ARNT. As additional controls, we confirmed the presence of the MOP1 and MOP2 proteins in the complex by showing that antisera raised against these proteins retarded the mobility of the complex.

To assay MOP1-ARNT and MOP2-ARNT interactions in vivo, we constructed a luciferase reporter driven by the hypoxia responsive TACGTG containing enhancer from the human EPO gene. Our transient expression experiments in Hep3B cells consisted of cotransfection of this reporter with vector control, MOP1, or MOP2 in the presence or absence of cobalt chloride to stimulate the hypoxia heme sensor. ARNT has been shown previously to be expressed in Hep3B cells. This experiment confirmed that the TACGTGcontaining enhancer sequence is responsive to cobalt and cotransfected MOP1 or MOP2 under normal oxygen tension. The transfected MOP1 construct appeared to be responsive to hypoxia (3.5-fold over control), while the MOP2 construct was only slightly responsive (1.2-fold). was more potent than MOP1 in driving expression of this reporter gene both in the presence and absence of cobalt chloride. This difference in efficacy of the MOP1 and MOP2 reporter plasmid in Hep3B cells could be explained by three possibilities: (1) the relative potency of the MOP2 transactivation domain may be much greater than MOP1; (2) the relative expression of MOP2 may be greater in this transient expression system than MOP1; or (3) the MOP1 may be partially repressed in vivo, by HSP90, while MOP2 is

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not (see HSP90 discussion below). Given that our MOP2 antisera are not useful in western blots, we could not assess the relative expression or stability of the MOP1 and MOP2 clones in this system.

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MOP3 is a conditionally active bHLH-PAS protein. Data from a number of laboratories suggests that Hsp90 represses AHR activity by anchoring the receptor in the cytosol away from its nuclear dimeric partner ARNT. Ligand binding appears to weaken the Hsp90 association and induce a translocation of the Hsp90-AHR complex to the nucleus where dimerization with ARNT can occur.

Two lines of evidence suggest that MOP3, like the AHR, may be a conditionally active bHLH-PAS protein and that in the absence of an unidentified cognate ligand, might be repressed and unable to dimerize in vivo. First, MOP3 interacts with HSP90 even more efficiently than human AHR, suggesting that MOP3 may be functionally repressed or anchored in the cytosol like the AHR. Second, MOP3 interacts with AHR in the coimmunoprecipitation assay, but not in the yeast interaction trap. Similarly, the AHR interacts with ARNT in the coimmunoprecipitation assay, but interacts weakly, if at all, in the absence of ligand activation.

Alternative explanations for the different MOP3-AHR interaction results obtained from our *in vitro* and *in vivo* systems should also be considered. For example, the structure of MOP3 may be different than the AHR and ARNT, such that positioning of the LexA domain adjacent to the bHLH-PAS domain may sterically hinder dimerization surfaces within this protein or lead to improper subcellular localization or instability of the chimera. One example of the potential negative impact of context sensitivity in the two-hybrid system can be observed in Figure 3. The LexAAHR-ARNT interaction is 14.7 times more robust than the LexAARNT-AHR interaction. In addition, the LexAAHR-ARNT interaction is more responsive to the AHR ligand \$NF\$ than the LexAARNT-AHR combination (6.4-fold and

2.0-fold, respectively). This difference cannot be explained by the relative transactivation potencies of the transactivation domains of AHR and ARNT in yeast, and therefore must be the result of context sensitivity. A final consideration is that coimmunoprecipitations may be capable of detecting weak interactions that cannot be maintained at the low cellular concentrations of the various MOPs. Thus, the MOP3-AHR dimerization may be too weak to occur in vivo. In this regard, we have previously reported ARNT-ARNT homodimers that bind specific DNA enhancer sequences in vitro, but they are weakly active, if active at all, in vivo (Swanson et al., 1995, supra).

It is also important to note that MOP1 and MOP4 also interact with HSP90 in the coimmunoprecipitation assay, albeit less strongly than MOP3 or human AHR. relatively weak interaction of MOP1 with HSP90 may be an indication that this protein is partially repressed in vivo and that it may have both constitutive and conditional activity. Such a phenomenon might explain why MOP1 has less transcriptional activity in our in vivo systems than MOP2, which does not interact with HSP90. Finally, MOP4 did not interact with the AHR or ARNT in either the coimmunoprecipitation assay or the interaction trap. Although our experience with AHR indicates that interactions with conditional bHLH-PAS proteins can be observed by coimmunoprecipitation assays, MOP4's interaction with HSP90 may also indicate a requirement for activation and may inhibit the sensitivity of detecting interactions in vivo.

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EXAMPLE 2 MOP3 Forms Transcriptionally Active Complexes with Circadian and Hypoxia Factors

As described above, a number of "orphan" bHLH-PAS proteins have emerged from searches of expressed sequence tag databases and low stringency hybridization screens. For newly discovered bHLH-PAS proteins that have close homologs (e.g., HIF1α and HIF2α (MOP2), or ARNT and

ARNT2), partnering and DNA binding specificity can often be predicted from amino acid sequence similarities in their bHLH-PAS domains. For divergent orphans like MOP3. MOP4, and MOP5, amino acid sequence does not provide the information necessary for similar predictions. To 5 characterize this class of orphans, we have employed a series of assays that allow us to: (i) identify heterodimeric partnerships, (ii) determine the DNA response element bound by these heterodimers, (iii) verify 10 that these complexes drive transcription in mammalian cells, and (iv) identify those tissues where these partnerships may occur. This example describes application of this approach to two bHLH-PAS orphans, MOP3 and MOP4.

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MATERIALS AND METHODS

Reagents. Oligonucleotides were supplied by GIBCO/BRL and designated as follows:

OL522 5'-GACAGTATCACGCCTCTCCTT-3'

20 OL579 5'-AGCGGCGTCGGGATAAAATGA-3'
OL595 5'-ATGCTGAACTGTGCCGAAAACTGT-3'
OL656 5'-GAACAGTGGGGTGCGTCCTCTTT-3'
OL990 5'-GGAATTCTGAGTCTGAAC-3'
OL991 5'-GGAATTCCACGCTCAGG-3'

OL992 5'-GGAATTCTGAGTCTGAAC(N)₁₃CCTGAGCGTGGATTCC-3'
OL1116 5'-GATCGGACACGTGACCATTGGTCACGTGTCCATTGGACACGTGACC-3'
OL1117 5'-GATCGGTCACGTGTCCAATGGACACGTGACCAATGGTCACGTGTCC-3'
OL1155 5'-GATCGGATACGTGACCATTGGTTACGTGTCCATTGGATACGTGACC-3'
OL1156 5'-GATCGOTCACGTATCCAATGGACACGTAACCAATGGTCACGTATCC-3'.

The yeast LexA fusion plasmid pBTM116 was provided by P. Bartel and S. Fields (State University of New York, Stony Brook). The yeast strain L40 was a kind gift of S. Hollenberg (Fred Hutchinson Cancer Research Center,

- Seattle, WA). The yeast strain AMR70 was constructed by Rolf Sternglanz, and was a kind gift of S. Hollenberg.

 LexA antiserum was a kind gift of J. W. Little

 (University of Arizona). pSGBCU11 was a kind gift of Stephen Goff (CIBA-Geigy, Research Triangle Park, NC).
- 40 Human CLOCK was a kind gift of T. Nagase (Kazusa DNA Research Institute, Chiba, Japan). Mammalian expression

vectors were purchased from GIBCO/BRL (pSVSport) and Promega (pTarget). Antibodies specific for MOP3 and MOP4 were prepared against peptides specific for each protein as described (Poland et al., 1991, Mol. Pharmacol. 39:20-26). The MOP3 peptide sequence was DNDQGSSSPSNDEAAC, and the MOP4 peptide sequence was KDKGSSLEPRQHFNALDVGC.

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Expression Plasmid Construction. expression plasmids harboring the LexA DNA binding domain fused to the bHLH-PAS domains of HIFl α (PL856), HIF2 α 10 (PL857), MOP4 (PL859), AHR (PL739), and ARNT (PL701) have been described (Example 1). LexAbHLH-PAS fusion plasmids for MOP3 (PL831) and CLOCK (PL828) were constructed in pBTM116 by an identical approach. Plasmids harboring the full-length ORFs of MOP3, MOP4, and CLOCK were constructed by PCR amplification of the ORF of each cDNA 15 and cloned into the appropriate vectors for expression in yeast or mammalian systems. For yeast expression of full-length proteins, PCR products were cloned into the appropriate sites of pSGBCU11. For mammalian expression, 20 PCR products were cloned into pSVSport and pTarget. yeast expression vector for full-length ARNT has been described (PL574) (Example 1). The yeast expression vector for full-length MOP3 was designated PL694. Mammalian expression vectors for ARNT (PL87), $HIF1\alpha$ 25 (PL611), and HIF2 α (PL447) have been described (Example 1; Jain et al., 1994, J. Biol. Chen 269:31518-31524). Mammalian expression vectors were constructed for MOP3 (PL691 and PL861), MOP4 (PL695 and PL871), and CLOCK (PL941).

Two-Hybrid cDNA Library Screen. The yeast interaction trap was performed using the yeast strain L40 (MATa, his3Δ200, trp1-901, leu2-3, 112, ade2, LYS::lexAop4HIS, URA3::lexAop8lacZ) or AMR70 (MATα, his3, lys2, trp1, leu2, URA::lexAop8-lacZ) as described (Example 1; Carver and Bradford, 1997, J. Biol. Chem. 272:11452-11456; Vojtek et al., 1993, Cell 74:205-214). The bait plasmid (PL859) was a fusion of the bHLH-PAS domain of

MOP4 to the DNA binding domain of LexA (Hogenesch et al., 1997, supra). The MOP4 bait construct was used to screen a human fetal brain cDNA library fused to the transactivation domain of Gal4 (CLONTECH) and 5 transformants were plated on selective media (minus tryptophan, uracil, histidine, and leucine). The cDNAs from surviving colonies, positive for lacZ activity were sequenced by the chain termination method (Sanger et al., 1977, PNAS 74:5463-5437). These sequences were analyzed using the BLAST algorithm (Altschul et al., 1990, J. Mol. Biol. 215:403-410).

Interaction Screen Against Known bHLH-PAS Proteins. LexAbHLH-PAS fusion proteins ("baits") of HIF1α, HIF2α, MOP3, MOP4, AHR, ARNT, and CLOCK were transformed into the L40 strain of yeast. The full-15 length ("fish") MOP3 and ARNT plasmids were transformed into the AMR70 yeast strain, and these transformants were plated onto yeast complete media plates (Kaiser et al., 1994, in Methods in Yeast Genetics, Cold Spring Harbor 20 Press, Plainview, NY). The L40 yeast harboring the bait constructs were replica plated onto these yeast complete media plates and mated for 8 hr at 30°C. The plates were then replica plated onto selective media and grown for an additional day at 30°C. 5-Bromo-4-chloro-3-indolyl 13-D-25 galactoside (X-Gal) overlay assays were performed to determine the relative expression of the lacZ reporter gene (Bohen and Yamamoto, 1993, PNAS 90:11424-11428). Western blot analysis, using LexA-specific sera, was performed on extract from each transformant to confirm 30 expression of the fusion protein (see Example 1).

DNA Binding Specificity. To determine high-affinity DNA binding sites for MOP3-MOP4 heterodimers, site selection and amplification was performed as described (Swanson et al., 1995, J. Biol. Chem. 270:26292-26302). Briefly, reticulocyte lysate expressed MOP3 and MOP4 proteins (~0.5 fmol each) were incubated with DNA oligonucleotide randomers corresponding to ~7

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X10⁷ different nucleotide sequences. Randomers were generated and amplified by PCR using oligonucleotides OL990 and OL991 as primers and OL992 as template. After incubating the complexes with the randomers for 30 min at 30°C, samples were loaded directly on 4% polyacrylamide—TBE (90 mM Tris/64.6 mM boric acid/2.5 mM EDTA, pH 8.0) gels to separate MOP3-MOP4 bound DNA from free DNA (Swanson et al., 1995, supra). Gel slices corresponding to the migration of bound DNA were excised, incubated overnight in TE (10 mM Tris/1 mM EDTA, pH 8.0), and the eluate subjected to additional PCR using oligonucleotides OL990 and OL991.

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Cell Culture and Transient Transfection.

Transient transfections of COS-1 cells were performed by the Lipofectamine protocol (GIBCO) as described in 15 Example 1. To mimic hypoxia, 100 μM of cobalt chloride was included in the cell growth media and incubated at 37°C until harvest. To monitor the transcriptional activity of the MOP3-MOP4 or MOP3-CLOCK heterodimers, a 20 synthetic reporter was constructed by annealing phosphorylated oligonucleotides OL1116 and OL1117 and cloning them into the BglII site in the reporter plasmid pGL3p (Promega). To measure the transcriptional activity of the MOP3-HIF1α or MOP3-HIF2α heterodimers, a synthetic 25 reporter was constructed by annealing phosphorylated oligonucleotides OL1155and OL1156 and then cloning them as above. Luciferase levels were reported in relation to β -galactosidase activity as described in Example 1.

antisense riboprobes, partial cDNAs of the mouse MOP3 and MOP4 were cloned into plasmid vectors harboring bacteriophage promoters. A partial 1.2-kb mouse fragment of MOP3 was obtained by PCR of a mouse kidney cDNA library using oligonucleotides OL579 and OL656, and cloned into pGEM-T in the T7 orientation. For MOP4, reverse transcription-PCR was performed on 3 µg of E17.5d placenta total RNA with oligonucleotides OL522 and OL595.

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The resultant fragment was subcloned in pGEM-T in the 5P6 orientation. Total RNA from various mouse tissues was prepared using the Trizol reagent (GIBCO/BRL) according to manufacturer's protocols. Ribonuclease protection assay (RPA) was performed as described for both MOP3 and MOP4 (Luo et al, 1997, Gene Expression 6:287-299). For in situ analysis, sense and antisense MOP3 and MOP4 riboprobes were generated with $[\alpha-[^{35}S]$ thio]UTP, 80 μ Ci (Amersham, >1,000 Ci/mmol; 1 Ci = 37 GBq) as the radioactive ribonucleotide and subjected to alkaline hydrolysis for 13 min at 60°C as described (Jain et al., 1998, Mech. Dev. 73:117-123). Tissue sections (5 μ m) were processed and hybridized with the specific riboprobes (Jain et al., 1998, supra).

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RESULTS

MOP4 Two-Hybrid Library Screen. The MOP4 bait plasmid was used to screen a human fetal brain cDNA library fused to the transactivation domain of Gal4. After screening 7 x 10⁵ colonies, 21 survived selection 20 and were blue in the presence of 5-bromo-4-chloro-3indoly1- β -D-galactoside. BLAST searches revealed that seven of these clones represented four independent MOP3 cDNA fragments. These cDNAs differed in their first 57 25 codons from the MOP3 cDNA we have described previously (GenBank accession no. U60415; SEQ ID NO:3). amino acids are identical to that reported by a second group, and appear to be derived from a second promoter (Ikeda and Nomura, 1997, Biochem. Biophys. Res. Commun. 233:258-264). All subsequent functional studies were 30 done using constructs derived from the MOP3 cDNAs identified by the yeast interaction trap.

MOP3 and MOP4 Screened Against Known bHLH-PAS
Proteins. To confirm the specificity of the MOP3-MOP4
interaction, we reversed the interaction trap strategy
and screened full-length MOP3 against all bHLH-PAS
proteins available in this laboratory. As a positive

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control we compared these results to a parallel screen using full-length ARNT. Western blot analysis using anti-LexA sera indicated approximately equal expression levels for all fusions. The full-length MOP3 protein interacted strongly with LexAbHLH-PAS fusions of MOP4, CLOCK, and HIF2\alpha and weakly with HIF1\alpha (Figure 5). No interaction of full-length MOP3 could be detected with LexA fusions of MOP3, AHR, ARNT, or the LexA control. Full-length ARNT demonstrated robust interactions with HIF2\alpha and the AHR, and weaker interactions with HIF1\alpha. We did not detect full-length ARNT interactions with LexAbHLH-PAS fusions of MOP3, MOP4, CLOCK, ARNT, or the LexA control (Figure 5).

DNA Binding Specificity of the MOP3-MOP4 15 Heterodimer. We performed a selection and amplification protocol to identify the DNA sequence bound with highaffinity by the MOP3-MOP4 complex. After three rounds of selection and amplification, a gel shift assay was performed using radiolabeled selected randomers to 20 identify the migration of the complex. We identified a species dependent on the presence of both proteins. A band corresponding to this migration was excised from the polyacrylamide gel, and used as template for a fourth round of amplification before cloning the pool. Analysis 25 of the sequencing data from 10 clones revealed that the MOP3-MOP4 heterodimeric pair bound the sequence G/TGA/GACACGTGACCC (Figure 6). This sequence is an imperfect palindrome containing a core E-box enhancer element (defined as CANNTG, underlined) and specificity for nucleotides in the flanking region (e.g., +4 "A"). 30 We refer to this response element bound by the MOP3-MOP4 as M34. To demonstrate sequence binding specificity and to confirm the selectivity for the +4 nucleotide, we performed competition experiment varying the +4 position 35 to A, C, G, or T (Figure 6). In agreement with our selection results, we observed a strong preference for the flanking +4 "A" nucleotide by the MOP3-MOP4 complex.

With MOP4 and CLOCK. To demonstrate that both MOP3 and MOP4 are required for binding to the M34 element, we performed additional gel shift experiments. A specific band was present only with the combination of MOP3 and MOP4, and was not present with either protein alone. As an additional specificity control, affinity-purified anti-MOP3 or anti-MOP4-specific Igs were used in gel shift experiments. Both MOP3-specific and MOP4-specific IgG were capable of retarding the mobility of the MOP3-MOP4 complex, while purified preimmune IgG alone was not.

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To determine whether the MOP3-MOP4 complex could drive transcription in vivo, we constructed a vector with three copies of the M34 element upstream of a minimal simian virus 40 promoter-luciferase reporter. Upon cotransfection of the reporter plasmid into COS-1 cells with MOP3 and MOP4, we observed that this combination enhanced transcription 3.3-fold, while neither protein alone was capable of driving transcription over control. The observations that CLOCK also interacted with MOP3 in the yeast interaction trap (Figure 5) and that CLOCK shares extensive homology with MOP4 prompted us to determine if MOP3-CLOCK complex could also drive transcription in vivo from an M34 element. Cotransfection of MOP3 and CLOCK revealed that this complex was also active, driving transcription 5.6-fold over control. Transfections with MOP3, MOP4, CLOCK, and ARNT alone, as well as combinations of ARNT and MOP3 or MOP4 failed to drive transcription over control.

MOP3 Forms Functional DNA Binding Complexes with HIFl α and HIF2 α . Prompted by our yeast interaction results, we set out to determine the ability of MOP3 to form DNA binding complexes with HIFl α in vitro. Because of the asymmetry at the +4 position of the M34 element, we were uncertain which half-site was bound by MOP3. Therefore, we synthesized enhancer elements with the HIFl α 5' half site (TAC) fused to both of the potential

MOP3 3' half-sites described above (GCCCTACGTGACCC or GCCCTACGTGTTCC). We found that the HIF1 α /MOP3 complex preferred the GCCCTACGTGACCC element in vitro, suggesting that MOP3 preferred an "A" at the +4 position. Therefore the corresponding response element bound by the HIF1 α -MOP3 complex, which we refer to as M13, was used in subsequent experiments. The results demonstrate that the M13 element is bound in the presence of the MOP3-HIF1 α combination, but not by either protein alone. MOP3-specific and HIF1 α -specific antisera abolished this complex while preimmune IgG did not. For comparison we included ARNT in these experiments, and found that ARNT-HIF1 α band was more intense than the MOP3-HIF1 α complex when all proteins were used at equimolar concentrations.

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15 To determine if MOP3 formed a transcriptionally active complex with either HIFla or HIF2a in vivo, we constructed a synthetic reporter using six copies of the M13 element described above. The M13 reporter was upregulated when cotransfected with combinations of MOP3-20 HIF1 α and MOP3-HIF2 α (3.3-fold and 3.6-fold, respectively). ARNT formed more active complexes with both HIFl α and HIF2 α (14.1-fold and 8.1-fold, respectively), consistent with our in vitro results. Like ARNT, upon exposure of these transfected cells to 25 cobalt chloride to simulate cellular hypoxia, MOP3 interacted and drove transcription in complexes with both $HIFl\alpha$ and with $HIF2\alpha$.

Coexpression of MOP3, MOP4, and HIF1a in Neonatal and Adult Murine Tissues. To determine if MOP3 was coexpressed with MOP4 in any murine tissue, ribonuclease protection assays (RPA) and in situ hybridization analysis were performed. Parallel RPA analysis of neonatal and adult tissues indicated that MOP3 was most highly expressed in brain, thymus, and muscle. MOP4 showed highest expression in the brain. We performed in situ hybridization analysis on tissues where RPA data indicated overlap between MOP3 and MOP4, or MOP3

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and HIFla. Sense controls were negative in all tissues except eye, where the pigment of the retina gave a nonspecific signal. In transverse sections of E15.5 brain, we observed that both MOP3 and MOP4 showed their highest expression levels in the thalamus. In E15.5 eye, we were able to detect colocalization of MOP3 and HIFlq in the retina, but were unable to detect specific The results show that both MOP3 and labeling of MOP4. $HIFl\alpha$ are colocalized in the thymic cortex of postnatal animals. Prompted by the observation of others that the MOP4 mRNA is expressed at low levels in the colon, we assayed that target tissue and observed that MOP4 and HIFlα were coexpressed in postnatal colonic mucosa, while MOP3 was undetectable there (Zhou et al., 1997, PNAS 94:713-718).

DISCUSSION

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In an effort to determine the pairing rules of MOP3 and MOP4, we employed the yeast interaction trap to identify the bHLH-PAS partners of these orphans. initial experiment using a MOP4 bait construct to screen a brain cDNA library identified MOP3 as a partner. further experiments, we reversed this approach and used full-length MOP3 to detect interactions with other bHLH-PAS members. This analysis confirmed the MOP3-MOP4 interaction and also demonstrated that CLOCK, $HIFl\alpha$ and HIF2α were additional partners of MOP3. As demonstrated previously, ARNT interacted with the AHR, $HIF1\alpha$, and $HIF2\alpha$, but not with MOP4 or CLOCK. The fact that both MOP4 and CLOCK interacted with MOP3 was not surprising given their 75% amino acid sequence identity in their bHLH-PAS domains. The observation that MOP3 was a partner of both $HIFl\alpha$ and $HIF2\alpha$, but that it did not dimerize with the AHR in the yeast interaction trap was an unexpected result. Due to lack of expression in our yeast system, we were unable to examine the interactions of MOP3 or MOP4 with a number of additional bHLH-PAS

proteins, including mSIM1, mSIM2, hARNT2, and hSRC1. Thus, we do not exclude the possibility that additional MOP3 and MOP4 interactions with these proteins may be important. Nevertheless, our data lead us to suggest that MOP3 is a general partner of a number of bHLH-PAS factors, with a distinct interaction profile from that of the more well characterized general partner ARNT.

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Analysis of MOP3 and MOP4 revealed that these proteins did not share perfect identity with any other known bHLH proteins in their basic residues thought to contact DNA. Therefore, we could not readily predict the response elements that the MOP3-MOP4 heterodimer would To overcome this limitation, we employed a DNA selection and amplification protocol and determined that the MOP3-MOP4 complex bound an E-box, with flanking region specificity for an "A" at +4 (i.e., CACGTGA, M34 element). In keeping with our prediction that MOP4 and CLOCK are functional homologs, transfection experiments demonstrated that the combination of either MOP3-MOP4 or MOP3-CLOCK was capable of driving transcription from M34 elements, while neither MOP3, MOP4, or CLOCK alone displayed this activity. In support of our argument that MOP3 harbors a partnering specificity distinct from that of ARNT, we observed that neither MOP3 nor MOP4 was capable of interacting with ARNT and driving transcription from the M34 element in its presence.

What could be the consequence of these interactions? Experiments from a number of laboratories indicate that circadian behavior may be regulated at the transcriptional level by complex interactions between multiple PAS domain containing proteins. Strong genetic evidence supports a role for CLOCK in the maintenance of circadian behavior in mice and the product of the period gene (PER) for control of circadian rhythms in Drosophila. The fact that MOP4 is a brain specific homolog of CLOCK and that these factors share MOP3 as a common dimeric partner suggests that both MOP3 and MOP4

may play a role in this process as well. In addition to the mammalian MOP3, MOP4 and CLOCK proteins, murine and human homologs of Drosophila PER have recently been characterized. Like Drosophila PER, the mRNA levels of these mammalian homologs respond to light and display Sequence analysis of PER proteins circadian rhythmicity. indicates that they contain PAS domains, but do not contain consensus bHLH domains. Coupled with additional biochemical evidence from others, these data suggest that PER proteins may affect their own transcription through interactions mediated by their PAS domains. Thus, these PERs may impact transcriptional activity of other bHLH-PAS dimers by acting as either dominant negative inhibitors that block pairing of trancriptionally active complexes, or they may act in a positive manner as coactivators of these complexes.

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In addition to defining the pairing rules and DNA binding specificities of MOP3 and MOP4, our data lead us to a testable model that describes circadian 20 oscillation of transcription. Without intending to limit the present invention by any particular explanation of mechanism, we speculate that MOP3-CLOCK or MOP3-MOP4 complexes regulate PER transcription through CACGTGAcontaining enhancers. The transcriptional activity of 25 these promoters could in turn be modified by feedback inhibition/activation by the PER protein products themselves. In support of this idea, an E-box element in the Drosophila PER promoter, required for normal cycling of the PER mRNA, bears striking resemblance to the M34 element we have identified (i.e., 5'-CACGTGAGC-3' 30 compared with 5'-CACGTGACC-3'). Given that we are borrowing from both Drosophila and mammalian systems, our model assumes that these signal transduction pathways have been largely conserved throughout evolution. keeping with this idea, a search of Drosophila expressed 35 sequence tags revealed the existence of an uncharacterized MOP4/CLOCK homolog (GenBank accession no.

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AA698290) and an uncharacterized MOP3 homolog (GenBank accession no. AA695336).

What could be the consequences of MOP3-HIF interactions? Transient transfection experiments showed that MOP3 formed transcriptionally active complexes with $\operatorname{HIFl}\alpha$ and $\operatorname{HIF2}\alpha$ and that these complexes responded to cellular hypoxia. MOP3 may play a specialized role in hypoxia signaling. The different tissue specific expression profiles of MOP3 and ARNT suggests that MOP3 may regulate cellular responses to hypoxia at distinct sites, such as the retina, thymic cortex, and thalamus. Moreover, the observation that MOP3 binds a GTG half-site with flanking region specificity for an "A" at +4, may indicate that MOP3/HIF complexes may have greater affinity for a distinct subset of hypoxia response elements (i.e., TACGTGA vs. the more commonly observed TACGTGG elements observed in known hypoxia responsive enhancers). Finally, the observation that MOP4 is expressed at a site where MOP3 expression appears low, i.e., colonic mucosa, suggests that additional partners may exist for MOP4 and CLOCK and that all bHLH-PAS signaling pathways may involve complex equilibria between multiple PAS proteins.

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EXAMPLE 3 Chromosomal Localization and Molecular Characterization of MOP7 as a Third Hypoxia Inducible Factor

Hypoxia inducible factors (HIFs) regulate transcriptional responses to low oxygen tension and other physiological conditions that rely upon glucose for cellular ATP. The HIFs are heterodimeric transcription factors that are composed of two bHLH-PAS proteins. The bHLH-PAS subunits can be classified as α -class or β -class. In addition to amino acid sequence similarity, the most distinguishing characteristic of the α -class subunits is that they are rapidly up-regulated by cellular hypoxia, or treatment with iron chelators and

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certain divalent cations (e.g. Co''). The previously described α -class subunits are referred to as HIF1 α (MOP1 herein) and HIF2 α (MOP2 herein). In contrast, the β -subunits appear to be constitutively expressed and ready to pair with their up-regulated α -class partner. Recent evidence suggests that ARNT, ARNT2 and MOP3 are prototype β -class subunits. At the present time, a number of well-characterized HIF-responsive gene products have been identified. These genes include those encoding EPO, VEGF and GLUT1, among others. The promoters of these genes are regulated by HRE elements that are recognized by the HIF $\alpha\beta$ heterodimer. The HREs contain the core TACGTG element and are found both 5' and 3' to the regulated promoter in a number of hypoxia responsive genes.

It is of academic and practical interest to 15 understand how bHLH-PAS proteins signal, as well as the biological consequences that result from the sharing of bHLH-PAS partners. The recent generation of thousands of expressed sequence tags (ESTs) has provided the opportunity to identify and classify orphan HIF subunits 20 based upon nucleotide sequence similarity with known bHLH-PAS proteins. As the result of these efforts, we have identified, and describe the cloning and characterization below, of a third HIF α -class subunit, 25 referred to above as MOP7. For consistency of nomenclature, this protein also is referred to as "HIF3 α ". Using a number of biochemical approaches, we demonstrate that the MOP 7 (HIF3a) cDNA encodes a protein that meets the major criteria of an α -class HIF subunit. The observation that multiple HIF α and β subunits are 30 encoded by the mammalian genome suggests that a complex array of subunit interactions and tightly controlled developmental expression patterns governs transcriptional response to hypoxic stress.

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MATERIAL AND METHODS

Gel-shift oligonucleotides. The complementary oligonucleotide pairs used in gel-shift assays are shown below (5' to 3'). They contain a constant flanking sequence and the wildtype or mutant HRE core sequence (underlined):

OL396 TCGAGCTGGGCAGGTAAGGTGGCAAGGC
OL397 TCGAGCCTTGCCACGTTACCTGCCCAGC
OL398 TCGAGCTTGCCAGGTGAGGTGGCAAGGC
OL399 TCGAGCCTTGCCACGTCACCTGCCAGC
OL414 TCGAGCTGGCAGGGTAGGTGGCAAGGC
OL415 TCGAGCCTTGCCACGTACCCTGCCCAGC

15 PCR Oligonucletides:

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OL1014
                   GCCATGGCGTTGGGGTGCAG
      OL1017
                   ACTGTGTCCAATGAGCTCCAG
      OL1178
                   GCCTCCATCATGCGCCTCACAATCAGC
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      OL1210
                   CCCCGTTACTGCCTGGCCCTTGCTCA
      OL1323
                   AGCCGAGGGGTCTGCGAGTATGTTGC
      OL1324
                   GCTGCTGACCCTCGCCGTTTCTGTAGT
      OL1397
                   GTCGACGCCACCATGGACTGGGACCAAGACAGG
      OL1427
                  GGATCCTCAGTGGGTCTGGCCCAAGCC
25
      OL1548
                   GCGGGGTGCTGGGAGTGGCTGCTAC
                  GCCTTCCTGCACCCGCCTTCCCTGAG
      OL1698
      OL1769
                  GCGGCCGCAAAAAACAAGACCGTGGAGACA
                  GCCCTGGGAGAATAGCTGTTGGACTTTGGGCAATTGCTCACT
      OL1771
      OL1772
                  GCGGCCGCCTATTCTGAAAAGGGGGGAAA
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      AP1
                   CCATCCTAATACGACTCACTATAGGGC
      AP2
                  ACTCACTATAGGGCTCGAGCGGC
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Cloning of MOP7. TBLASTN and BLASTX algorithms 35 were used to search nucleotide sequences corresponding to amino acids 54 to 125 of hHIF1 α (http://dot.imgen. bcm.tmc.edu: 9331/seq-search/Options/blast.html) (Hwang et al., J. Mol. Cell. Cardiol. 26: 1329-1333, 1994). mouse EST clone, Genbank Accession AA028416 (designated 40 PL773), was found to encode a novel bHLH-PAS protein. obtain the complete open reading frame, we performed a series of PCR amplifications using primer-anchored cDNA derived from mouse lung ("Marathon-Ready," Clontech) (Siebert et al., Nuc. Acids Res. 23: 1087-1088, 1995). A 45 3' rapid amplification of cDNA ends (RACE) reaction was performed using oligonucleotides OL1178 and anchor primer The product of this reaction was reamplified in a second reaction with OL1178 and AP2. The 2.0-kb 3' PCR product obtained by this protocol was cloned into the

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pGEM Teasy vector (Promega) and designated PL970. clone was sequenced and found to contain an ORF followed by a translational stop site (Figure 7). To confirm the position of the traslational stop site, OL1324 was used in an independent 3' RACE reaction. The 0.9 kb product was cloned into a pGEM Teasy vector (PL1017) and was found to contain the same stop codon (Figure 7). To obtain the 5' end of the cDNA, OL1323 was used in a RACE reaction against oligonucleotide AP1. The 1.2 kb RACE product was cloned into pGEM Teasy vector (PL1016) and 10 found to contain a translation start codon ATG followed by a long ORF. We define the nucleotide A from the initiation codon as position 1 of the cDNA. In addition, the translational start site is defined by the presence of an in-frame stop codon 51 nucleotides upstream. generate expression plasmids containing the full ORF, a PCR reaction was performed using OL1210 and OL1397 with PL1016 as template. The PCR fragment was cloned into pGEM Teasy vector in the SP6 orientation and named PL1024. The NdeI digested PL1024 was then inserted into the NdeI digested PL970 to generate the full-length HIF 3α in the pGEM-Teasy vector (PL1025).

Construction of MOP7 expression plasmids. MOP7 expression in mammalian cells, the ORF was amplified by PCR using OL1397 and OL1427 with PL1025 as template. The resultant plasmid was cloned into pTarget vector downstream of the CMV promoter (Promega) and was named PL1026 (Figure 7).

To confirm the hypoxia inducibility of MOP7, we constructed a fusion protein comprised of the DNA binding domain from GAL4 (residues 1-147), the predicted hypoxia responsive domain-1 (HRD1) from mMOP7 (residues 453-496), and the transactivation domain (TAD) from hARNT (residues 581-789). The HRD1 was amplified using OL1769 and OL1771 with mMOP7 as template. To form the HRD1/TAD chimera, the resultant PCR fragment from above was used as a megaprimer in a second PCR reaction with OL1772 as the

second primer and hARNT as the template (Barik et al., Biotechniques <u>10</u>: 489-490, 1991). The HRD1/TAD chimeric fragm nt was cloned into the *Not*I site of the GAL4 fusion vector pBIND (Promega) and designated PL1131.

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Structural gene analysis and chromosomal The MOP7 insert from PL773 was cut with localization. EcoRI/NdeI and the 0.6-kb fragment was purified and used as a probe to screen for bacterial artificial chromosome (BAC) clones containing the mouse MOP7 gene (Genome Systems, Inc.). Oligonucleotides derived from the mMOP7 sequence were used as primers to sequence the BAC DNA, and the splice sites were deduced by comparing the genomic and cDNA sequences. To obtain BACs containing the human MOP7, oligonucleotides OL1014 and OL1017 were used in a PCR reaction with human heart cDNA as template (Clontech) to amplify a MOP7 fragment (Genbank Accession No. AF079154). This fragment was subcloned into the pGEM-Teasy vector, confirmed by sequencing, and used as a probe to screen for a BAC clone harboring the human structural gene for MOP7 as above. The identity of the resultant BAC was confirmed by direct sequencing using primers specific for hMOP7. The human MOP7 chromosomal location was identified by PCR reactions against human/hamster somatic cell hybrid DNA using human MOP7specific oligonucleotides. This location was confirmed by fluorescence in situ hybridization (FISH) using the BAC harboring human MOP7 structural gene as the probe (Genome Systems, Inc.).

Northern Blot analysis. To generate a hybridization probe for northern blot analysis, the 894 bp MOP7 insert from PL1017 was excised with EcoR1 and radiolabeled with $[\alpha^{-32}P]dCTP$ by random priming. A northern blot containing poly A' mRNA from different mouse tissues (Origene Technologies, Rockville, MD) was hybridized with 5 x 10^6 cpm/ml MOP7 probe. β -actin was used as a loading control.

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Gel-shift assay. To generate a double strand oligonucleotide probe containing the core HRE element, 50 ng of oligonucleotide OL414 was end-labeled with [y-³²P|ATP and was annealed with 10 fold excess of cold complementary oligonucleotide OL415. Unlabeled oligonucleotides containing either wild-type HRE sequence (TACGTG) or mutant HRE sequences, AACGTG (OL396/397) or GACGTG (OL398/399), were used in competition experiments to demonstrate specificity. For expression of the bHLH-PAS proteins, mMOP7 (PL1025) and hARNT (PL87) were synthesized in a reticulocyte lysate in the presence of [35S]methionine. The amount of each protein synthesized was calculated by measuring radioactivity and estimated to be approximately 1 fmol in each 10 μ l gel-shift reaction. Each gel-shift assay was performed with 100,000 cpm of oligonucleotide probe per 10 μ l reaction. To confirm complex identity, 1 μ l of anti-ARNT sera was used to supershift the DNA bound protein complex.

Cell culture and transfection. COS-1 cells were maintained in high glucose DMEM medium supplemented 20 with 10% fetal calf serum, 100 units/ml penicillin and streptomycin. The HRE driven luciferase reporter (PL945) was made by annealing OL1174 and OL1175 and then cloning the fragment into pGL3-promoter vector (Promega, Madison, 25 WI). For transient transfection experiments, mammalian expression plasmids expressing MOP7 or hARNT with the reporter using lipofectamine (GIBCO BRL Life Technologies). A β -galactosidase-expression plasmid was co-transfected to control for transfection efficiency. Cells were incubated for 20-24 hours with or without 30 treatment of cobalt chloride or hypoxia (1% 02) before being harvested. The luciferase and β -galactosidase activities were determined using the luciferase assay and the Galacto-Light protocols as previously described (see 35 Example 2).

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RESULTS and DISCUSSION

From a TBLASTN search of the dBEST database with the sequence corresponding to amino acid residues 54 to 125 of hHIF1\alpha, we identified a mouse EST clone (AA028416) that appeared to be a novel bHLH-PAS protein. This protein is referred to herein as MOP7 and as $HIF3\alpha$. to denote its relationship to other hypoxia-inducible factors. To obtain the complete ORF of this cDNA, a series of RACE reactions was performed using cDNA from mouse lung as template. The MOP7 ORF (SEQ ID NO:7) spans 1.98 kb and encodes a 662-amino acid protein (SEQ ID NO:16) with a predicted molecular weight of 73 kDa. Northern blot analysis on mRNA prepared from selected mouse tissues identified a MOP7 transcript that is expressed in adult thymus, lung, brain, heart and kidney. This expression pattern is distinct from that reported for other α -class HIFs. HIF1 α is most abundantly expressed in kidney and heart, and $HIF2\alpha$ is most abundantly expressed in vascular endothelial cells and is highest in lung, placenta and heart.

HIF1 α (MOP1) is the most well-characterized α class subunit. This protein can be described by a number of signature motifs. In addition to the bHLH-PAS domains, HIF1 α also contains two HRD motifs in its Cterminus that confer hypoxia responsiveness. appears to primarily confer hypoxia-dependent protein stability, whereas HRD2 appears to confer hypoxiaresponsive transcriptional activity. To determine if similar motifs occur in MOP7, we compared HIF1\alpha, HIF2\alpha and MOP7 protein sequences using the CLUSTAL algorithm (Higgins & Sharp, Gene 73: 237-244, 1988. These three sequences shared greater than 92% identity in the basic region, 68% in the HLH domain, and greater than 53% in the PAS domain. Although little sequence with significant homology to HRD2 was found, a 36-amino acid stretch within the C-terminal half of MOP7 was found to share 61% identity with the HRD1 of HIF1 α .

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To further demonstrate the evolutionary relationship between these α-class HIFs, we compared their gene structure and chromosomal localization. Direct sequencing of a BAC clone containing the mMOP7 gene revealed 15 exons, all with consensus splice donor/acceptor sites (see sequences of Genbank Accession No. AF079140-079153 for exon-intron junctions). We found that 11 of 15 and 10 of 15 splice junctions found in the mMOP7 gene are conserved to those found in hte structural genes of mHIF1 α and hHIF2 α , respectively (Figure 8). characterize the distribution of HIF genes in the mammalian genome, we used human MOP7-specific PCR reactions against a human/hamster somatic cell panel and mapped the MOP7 gene locus on human chromosome 19. locus was further defined to chromosome 19q13.13-13.2 by FISH using a BAC clone containing the human MOP7 structural gene as a probe. Therefore, the human MOP7 locus is distinct from that of human HIF1\alpha and HIF2\alpha. which reside on chromosome 14q21-24 and 2p16-21, respectively.

As a biochemical proof that MOP7 was a bona fide α-class HIF, we performed gel-shift and transient transfection analyses. Because HIF1\alpha and HIF2\alpha are known to pair with the β-class HIF subunit ARNT, we predicted that MOP7 would also pair with ARNT. Based upon sequence identity in their basic regions, we also predicted that a MOP7-ARNT would bind the HRE core sequence, TACGTG. predicted, the gel-shift analysis showed that MOP7 only bound to the HRE containing oligonucleotide in the presence of ARNT. The specificity of the interaction was demonstrated by two additional observations. First, the MOP7-ARNT-HRE complex was abolished by anti-ARNT IgGs but not by preimmune antibodies. Second, the complex was blocked by an excess of HRE containing oligonucleotide but not by oligonucleotides with a single mutation within the core HRE sequence. To determine if this interaction could also occur in vivo, MOP7 and/or ARNT were

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cotransfected into COS-1 cells with a luciferase reporter driven by six HRE enhancer elements. The results demonstrated that the combination of MOP7 and ARNT upregulated transcription from the HRE-driven reporter by 11.7 fold, whereas neither protein alone had an effect. In addition, the activity of these complexes was enhanced by either hypoxia or cobalt chloride.

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To demonstrate that the MOP7 activity was directly upregulated by hypoxia, we employed a fusion protein approach that has been used to map the HRDs of HIF1α. HRD1 of HIF1α has been shown to encode a hypoxiaresponsive protein stability domain that also displays weak transcriptional activity. Given the sequence similarity between residues 453-496 if MOP7 and the HRD1 of HIFla, we predicted that this domain would independently respond to hypoxic stimulus or cobalt ion To test this, we constructed a plasmid expressing a fusion protein comprised of the DNA binding domain of GAL4, the predicted HRD1 of MOP7, and the TAD from ARNT. We predicted that we could measure the response of this domain by monitoring the output from a GAL4-driven luciferase reporter in Hep3B cells. results demonstrated that the fusion protein's activity increased by 4.5- and 4.2-fold, upon treatment with cobalt chloride or hypoxia, respectively. In control experiments, we observed that a GAL4 fusion protein harboring only the ARNT-TAD did not respond to either hypoxia or cobalt chloride treatment. The level of inducibility seen with the HRD1 fusion is consistent with that obtained for a similar fusion protein using the HRD1 domain of HIF1\alpha. This result provided evidence that amino acids 453 to 496 of MOP7 was sufficient to confer the hypoxia inducibility and that the stability of the parent protein is regulated in a manner that is similar to that of HIF1 α and HIF2 α .

In eucaryotes, transcriptional responses to low oxygen tension are mediated by complex interactions

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between a number of α - and β -class HIF subunits. characterization of a third α-class HIF with a tissue distribution that is distinct from either HIF1 α or HIF2 α provides evidence that cellular responses to hypoxia result from a complex set of interactions from multiple combinations of $\alpha\beta$ pairs. MOP7 also may have a distinct role in mediating biological responses to hypoxia. support of this notion, MOP7 and HIF1α have limited sequence homology in their C-termini. Most importantly, 10 MOP7 contains sequence that corresponds to HIF1α's protein stability element, HRD1, but not to its hypoxiaresponsive TAD element, HRD2. This may indicate that MOP7-ARNT complexes have decreased transcriptional potency relative to other HIF heterodimers. 15 importance of this complexity is underscored by the presence of HIF1 α , HIF2 α and MOP7 in both mice and Finally, this complexity appears to be highly conserved among vertebrates. In support of this idea, we have cloned a partial human MOP7 cDNA and have shown all 20 three HIF \alpha-class genes reside on separate human chromosomes and display considerable sequence divergence in their C-termini.

The present invention is not limited to the
25 embodiments described and exemplified above, but is
capable of variation and modification within the scope of
the appended claims.

w claim:

- An isolated nucleic acid molecule that includes an open reading frame encoding a protein selected from the group consisting of: MOP2, MOP3, MOP4, MOP5, MOP6 MOP7, MOP8 and MOP9.
 - 2. The nucleic acid molecule of claim 1, which is DNA.

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- 3. The DNA molecule of claim 2, which is a gene, the exons of which comprise said open reading frame encoding said protein.
- 15 4. The DNA molecule of claim 2, which is a cDNA.
 - 5. The nucleic acid molecule of claim 1, wherein said open reading frame encodes a protein having an amino acid sequence substantially the same as a sequence selected from the group consisting of SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17 and SEQ ID NO:18.
- 6. The nucleic acid molecule of claim 5, which comprises a sequence substantially the same as a sequence selected from the group consisting of SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8 and SEQ ID NO:9.
- 7. An isolated protein, which is a product of expression of part or all of the open reading frame of claim

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- 8. A recombinant DNA molecule comprising the nucleic acid molecule of claim 1, operably linked to a vector for transforming cells.

- 9. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:2;
 - b) an allelic variant of SEQ ID NO:2:
 - c) a natural mutant of SEQ ID NO:2;
- d) a sequence specifically hybridizing with part or all of a sequence complementary to SEQ ID NO:2 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:2; and
- e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:11.
 - 10. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
- 15 a) SEQ ID NO:3;
 - b) an allelic variant of SEQ ID NO:3:
 - c) a natural mutant of SEQ ID NO:3;
 - d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:3 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:3; and
 - e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:12.
- 25 11. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:4;
 - b) an allelic variant of SEQ ID NO:4:
 - c) a natural mutant of SEQ ID NO:4;
- d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:4 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:4; and
- e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:13.

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- 12. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:5;
 - b) an allelic variant of SEQ ID NO:5;
 - c) a natural mutant of SEQ ID NO:5;
- d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:5 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:5; and
- e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:14.
 - 13. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:6;
 - b) an allelic variant of SEQ ID NO:6;
 - c) a natural mutant of SEQ ID NO:6;
 - d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:6 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:6 and
 - e.) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:15.
- 25 14. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:7;
 - b) an allelic variant of SEQ ID NO:7;
 - c) a natural mutant of SEQ ID NO:7;
- d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:7 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:7; and
- e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:16.

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- 15. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:8;
 - b) an allelic variant of SEQ ID NO:8;
 - c) a natural mutant of SEQ ID NO:8;
- d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:8 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:8; and
- 10 e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:17.
 - 16. An isolated nucleic acid molecule having a sequence selected from the group consisting of:
 - a) SEQ ID NO:9;
 - b) an allelic variant of SEQ ID NO:9;
 - c) a natural mutant of SEQ ID NO:9;
 - d) a sequence hybridizing with part or all of a sequence complementary to SEQ ID NO:9 and encoding a polypeptide substantially the same as part or all of a polypeptide encoded by SEQ ID NO:9; and
 - e) a sequence encoding part or all of a polypeptide having amino acid SEQ ID NO:17.
- 17. A recombinant DNA molecule comprising the
 25 nucleic acid molecule of any of claims 9-16, operably linked
 to a vector for transforming cells.
 - 18. An oligonucleotide between about 10 and about 100 nucleotides in length, which specifically hybridizes with a portion of the nucleic acid molecule of any of claims 9-16.
- 19. The oligonucleotide of claim 18, wherein said portion includes a translation initiation site of said polypeptide.

- 20. A cell transformed with the recombinant DNA molecule of any of claims 9-16.
- 21. An isolated protein encoded by the nucleic 5 acid molecule of any of claims 9-16.
 - 22. An antibody immunologically specific for part or all of the protein of claim 8.
- 10 23. An antibody immunologically specific for part or all of the protein of Claim 21.
- 24. An isolated nucleic acid molecule comprising a sequence that encodes an α -class hypoxia-inducible factor, said factor being encoded by a gene located on human chromosome 19q13.13-13.2.
- 25. A polypeptide encoded by the nucleic acid molecule of claim 24, which is produced in mammalian adult thymus, lung, brain, heart and kidney.
 - 26. The polypeptide of claim 25, having amino acid SEQ ID NO:16.
- 27. An isolated nucleic acid molecule comprising a sequence that encodes an α -class hypoxia-inducible factor, said factor being encoded by a gene located on human chromosome 2p16-21.
- 30 28. A polypeptide encoded by the nucleic acid molecule of claim 27, which is produced in mammalian vascular endothelial cells.
- 29. The polypeptide of claim 28, having amino 35 acid SEQ ID NO:11.

- 30. An isolated nucleic acid molecule comprising a sequence that encodes a circadian-responsive factor that heterodimerizes with a binding partner and stimulates transcription of genes having an E-box transcription control element comprising a sequence CACGTGA.
- 31. The nucleic acid molecule of claim 30, wherein said sequence encodes a protein having an amino acid sequence substantially the same as a sequence selected from the group consisting of SEQ ID NO:13 and SEQ ID NO:17.
- 32. The nucleic acid molecule of claim 30, which comprises a sequence substantially the same as a sequence selected from the group consisting of SEQ ID NO:4 and SEQ ID NO:8.
- 33. An isolated protein, which is a product of expression of part or all of the nucleic acid molecule of claim 30.

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- 34. A recombinant DNA molecule comprising the nucleic acid molecule of claim 30, operably linked to a vector for transforming cells.
- a sequence that encodes a binding partner for a circadianresponsive factor that heterodimerizes with said binding
 partner and stimulates transcription of genes having an Ebox transcription control element comprising a sequence
 CACGTGA.
 - 36. The nucleic acid molecule of claim 35, wherein said sequence encodes a protein having an amino acid sequence substantially the same a sequence selected from the group consisting of SEQ ID NO:12 and SEQ ID NO:18.

37. The nucleic acid molecule of claim 35, which comprises a sequence substantially the same a sequence selected from the group consisting of SEQ ID NO:3 and SEQ ID NO:9.

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- 38. An isolated protein, which is a product of expression of part or all of the nucleic acid molecule of claim 35.
- 10 39. A recombinant DNA molecule comprising the nucleic acid molecule of claim 35, operably linked to a vector for transforming cells.
- 40. A method for determining if a test compound regulates circadian function, which comprises the steps of:
 - a) providing a recombinant cell comprising:
 - i) a heterologous nucleic acid molecule encoding an expressible circadian-responsive factor that forms a heterodimer with a binding partner and stimulates transcription of genes having an E-box transcription control element comprising a sequence CACGTGA;
 - ii) a heterologous nucleic acid molecule encoding the binding partner for the circadian-responsive factor; and
- iii) a reporter gene comprising at least one transcriptional control element having a sequence CACGTGA, operably linked to a reporter coding sequence, the expression of said gene being induced by binding of said heterodimer to said transcriptional control element;
 - b) preparing a culture of the recombinant cells;
 - c) incorporating the test compound into the cell culture under conditions permitting expression of the heterologous nucleic acid molecules, heterodimerization of the factor and the binding partner, and binding of the heterodimer to the transcriptional control element;

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- e) measuring expression of the reporter gene and comparing the expression to an equivalent cell culture in which the test compound was not incorporated, a change in expression of the reporter gene in the culture containing the test compound being indicative of the ability of the test compound to regulate circadian function.
- 41. The method of claim 40, wherein the circadian-responsive factor is MOP4 and the binding partner 10 is MOP3.
 - 40. A method for determining if a test compound regulates red blood cell synthesis, glucose metabolism or angiogenesis, which comprises the steps of:
 - a) providing a recombinant cell comprising:
 - i) a heterologous nucleic acid molecule encoding an expressible α -class hypoxia-responsive factor that forms a heterodimer with a binding partner and stimulates transcription of genes having a hypoxia responsive transcription control element (HRE);
 - ii) a heterologous nucleic acid molecule encoding the binding partner for the hypoxia-responsive factor; and
- iii) a reporter gene comprising at least one transcriptional control element having a sequence CACGTGA, operably linked to a reporter coding sequence, the expression of said gene being induced by binding of said heterodimer to said HRE;
- b) preparing a culture of the recombinant cells;
 - c) incorporating the test compound into the cell culture under conditions permitting expression of the heterologous nucleic acid molecules, heterodimerization of the factor and the binding partner, and binding of the heterodimer to the HRE;

- e) measuring expression of the reporter gene and comparing the expression to an equivalent cell culture in which the test compound was not incorporated, a change in expression of the reporter gene in the culture containing the test compound being indicative of the ability of the test compound to regulate red blood cell formation, glucose metabolism or angiogenesis.
- 43. The method of claim 42, wherein the hypoxia10 responsive factor is selected from the group consisting of MOP1, MOP2, MOP6 and MOP7.

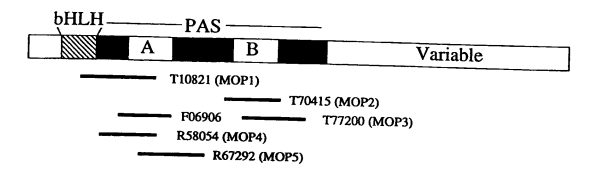


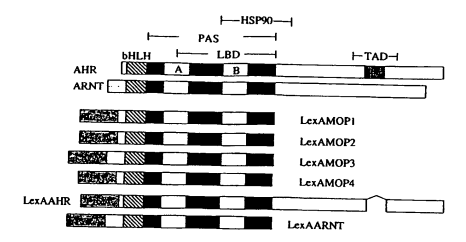
FIGURE 1

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IIDDOOLEEVPLYNDYNLP MOP1 FESSAYGKAILPPSOPWA MOP2 SSSPLNISTPPPDASSPG MOP3 STPIKLMAEASIPALPRSA MOP9 SPSSHPAIPRPEFISVIRA MOP5	FPNFESLELSFTMPOTODO MOPT TPNSPEDYTSLDNOLKIE MOP2 HIGIOMIONDOGSSSPRWO MOP3 TPAPMAOFSADFSHFOTIK MOP4 IRYGPAELGLYYPHLORLG MOP5	LFAEDTEAKNPFSTODTOL MOP1 OLSPICPEERLLAENPOST MOP2 MOP3 OPAVSLSFSSTORPEAGOD MOP4 MOP5	HOP HOP 1	1100 H 100 P 100 P 100 P	ACRLLGOSHDESGLPOLIS MOP1 PLPOPPSAISPGENSKSRF MOP2 RYDREVKVPVLGSSILLOG MOP2	нор2	
VESEDISSLFDKLKKEPDAL ILLAPAAGDIIISLDFGSNDIE IDDOOLEEVPLYNDVNLP GAVSEKSNFLFIKLKEEPEELAGLAPIPGDAIISLDFGNONFEESSAYGKAILPPSOPWA THFIVPGIPGGTRAAGKIGRHIAEEINEIMRINGSLRSSCSSPLNITSPPDASSPG EPROHFNALDVGASGLNISMSPSASSRSSHKSHIAMSEPISTPIKLMEASIPALPRSA PPIEGKQAAPAENEAPGIGGKRIKVEPGFREIKGSEDSGDEGPSSHPAIPRPEFISVIR	SPNEKLONINLAHSPLPIAETPKPLRSSADPALNOEVALKLEPNPESLELSFINPOIGDO TELSHSIDSEAGSLPAFIVPDAAPFGSI TPSATSSSSSCS IPNSFEDTYISLDNDLKIE GKKILNGITDIPSSGLLSGDAGENPGVPTSDSSSILGENPHIGIDHIDNDGSSSPSND GKKILNGELPPIPSSGLLSGDAGENPGVPTSDSSSILGENPHIGIDHIDNDGSSSPSND TLPDELLPPAFLSDAAFHAPPFSSCOLTOGLLPOIVLGS IPAPHAOFSAOFSHFOIIK GVLKODPVRPWGLAPPGDPPTILLHAGFLPPVVRGLCIPGIIRYGPAELGLVYPHLORLG	TPSPSDGSTROSSPEPNSPSETCFYVDSDHVNEFKLELVEKLFAEDTEAKNPFSTDDTDL VIEKLFAHDTEAKDGCSTOTDFNELDLETLAPYIPHDGEGFOLSPICPEERLAENPOST EAAKAVIHSLLEADAGLGGFYDFSDLPWPL DOLEGRIRILGANIRWOGEELHKIGEOLCLVDDSNYOHFLOOPAVSLSFSSTORPEADOO PGPALPEAFYPPLGLPYPGPAGTRLPRKGD•	OLEMLAPY IPHODDFOLRSFOOLSPLESSSASPESASPOSIVIVFOOIOIOEPTANAIII POHCFSAMINIFOPLAPVAPHSPFLLOKFOOOLESKKIEPERRPHSSIFFDAGSKASLPP LOORSAAVIOPOLGAGPOLPGOISSAOVISOHLLRESSVISIOGPKPHRSSOLMOSSGRS	CCGDASIPLSHGGRSNIOWPPDPPLHFGPIKWAVGDORIEFLGAAPLGPPVSPPHVSIF IPAERRSLPLFVGLVS EDIEKSHPRSPNVLSVALSORIIVPEEELNPKILALDNAGRKKHEHOGSLFGAVGIGTL KTRSAKGFGARGPNVLSVALSOKLIVVEKKKKROLEYEKOAFODPSGGDPFGGSISHLMWKR	LOOPDDHAATTSLSWKRYKGCKSSEONGHEOXTIILIPSDLACRLLGOSHDESGLPOLTS HKNLRGGSCPLHPDKPLSANVPNDKLTONSHROLGHPLRHLPLPOPPSATSPGENSKSRF YDCEVNAPIOGSRNLLGGEELLRALDOVN• PPOCYATOYODYSLSSAHKYSGAASRLLGPSFESYLLPELTRYDREVKVPYLGSSTLLOG	114.	
) GDLLRALDOAIH•	
378 475 388 332	438 440 635 363 363	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2880 2880 880 880	620 620 626 678 680	738 740 798 800	880	
HOP 2 HOP 3 HOP 4 HOP 5	1007 1007 1007 1007 1007 1007 1007 1007	700 700 700 700 700 700 700 700 700 700	MOP2 MOP2 MOP3 MOP4	HOP2 HOP2 HOP4 HOP5	100 H	HOP 1 HOP 3 HOP 4 HOP 5	HOP 1 HOP 2 HOP 4 HOP 4
I MEGAGGAMDKKKISSER 1 HIAOKEKKRSSER 1 HIAOKEKKRSSSER 1 HIAOKEAVSLWALIVSLOPPVPLCVCREHIGSGRRKOOCVILPFISRELCFYLLFPPPRLEYTEMGGGIKN 1 NDEDEKORA 1 NDEDEKORA	X X 4 X 1	GOLDIEDDHKAONNCFYLKALGFVWVLID SENESEAEADDOMDNLYLKALEGFJAVVID SENESEAE	128 GHSVFDT IMPCDHEEMERHITHR	, •	205 SNOPOCCYKRPPHICLYLICEPIPHPSNIEIPLOSKIFLSRHSLDHKFSYCDER 207 PHNSLCGYKEPLISCLINCEPIQHPSHHDIPLOSKIFLSRHSMOHKFIYCDOR 302 PONEGCNLSCLVAIGRLHSHVVPOPVNGE RVKSHEYSRNAIDGKFVFVDOR 223	259 TIELMGYEPEELLGRSIYEYYHALDSDHLIKIHHOHFIT-KGOYTIGOYRHLAKRGGYWWY 281 TIELIGYHPEELLGRSAYEFYHALDSENHIKSHONLGT-KGOYVSGOYRHLAKRGGYWWY 285 ATALLAYLPOELLGTSCYEYTHODOIGHLAECHRONLDTREKITINCYKFKIKOGSFTTL 287 APPIIGYLPFEVLGTSGYOYYHIDOLELLARCHONLMGF-GKGSCCYRFLIKGOOWIWL 213 LYSDHMOLGPSELYGFSYNGOAIRLROSHWOLLD-KGOYHIGYYRWLORAGGFWHL	318 EIDATY ITRIKNSOPOCITYVYYSGI 10HDL IFSLODIECYLRPVESSDHKHIOLF TK 320 EIDGTY IYNPRNLOPOCIHCVNYYLSE IEKNDYVFSHOOTESLFKPHHAHNSI FDSSGK 416 RSRWFSFHNPWIKE VET IVS INTVVLANVLEGGOP IF POL I ASPHISNDSHLPSGEGGPKR 328 OIHYYI ITHOWNSKPEF IVCTHSVYSYAOVRVERROELALEDPPSEALHSSALKOKGSSL 272 OSVAIYAGSGKSPGEHHVLWVSHVLSOAEGGOIPLDAFOLPASVACEEASSPGPEP IEPE
HSKEAVSL	158 7.5 29	76 75 131 34		± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ±	~%%% <u>~</u>	ฉฉฅฉีฉ	ઌઌૻ ૱ઌૢઌ
_							

FIGURE 2

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FIG. 3A



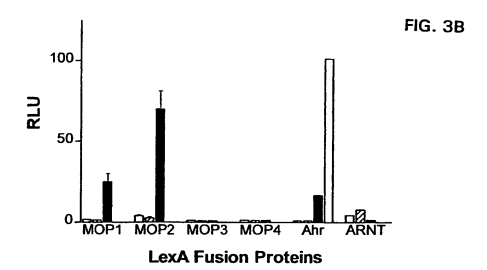


FIGURE 3

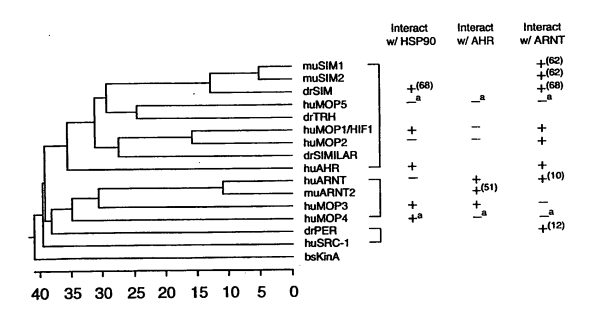
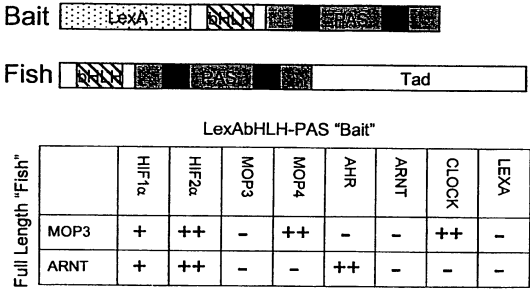


FIGURE 4

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g	9	G	G	C	Α	С	G	Ţ	G	Α	С	Α	С
G	G	T	Α	C	Α	С	G	T	G	Α	С	С	С
t	g	а	а	C	Α	C	G	T	G	Α	С	С	С
t	g	а	а	C	Α	С	G	T	G	Α	С	T	C
g	g	G	С	C	Α	C	G	T	G	Α	С	С	T
G	G	G	Α	C	Α	С	G	T	G	Α	С	С	g
С	T	Α	Α	C	Α	С	G	T	G	Α	С	С	Ğ
g	а	а	С	C	Α	C	G	T	G	Α	G	С	ī
t	g	а	а	C	Α	C	G	T	G	Α	С	Α	C
9	G	G	T	С	Α	С	G	T	G	Α	С	T	С
G/T	G	A/G	Α	С	Α	С	G	Т	G	Α	С	С	С
-7	-6	-5	-4	-3	-2	- 1	+1	+2	+3	+4	+5	+6	+7

FIGURE 5



HIF1α HIF2a MOP3 MOP4 CLOCK AHR R ARNT LEXA MOP3 + ++ ++ ++ **ARNT** + ++ ++

FIGURE 6

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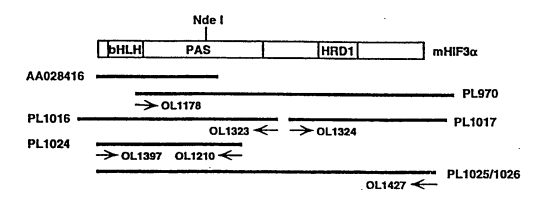


FIGURE 7

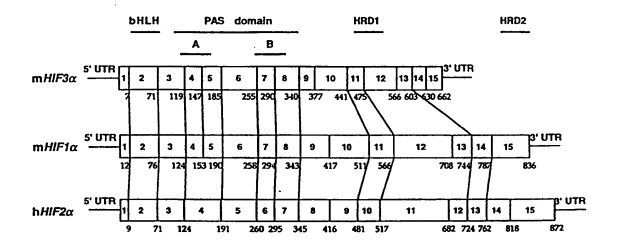


FIGURE 8

NUCLEOTIDE AND DEDUCED AMINO ACID SEQUENCES

MOP1:

Sequence I.D. No. 1

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1 cacgaggeag cactetette gtegettegg ccagtgtgte gggetgggee etgacaagee
  61 acctgaggag aggeteggag eegggeeegg acceeggega ttgeegeeeg etteteteta
121 gtctcacgag gggtttcccg cctcgcaccc ccacctctgg acttgccttt ccttcttc
181 tccgcgtgtg gagggagcca gcgcttaggc cggagcgagc ctggqqqccg cccqccqtqa
241 agacategeg gggacegatt caccatggag ggegeeggeg gegegaaega caagaaaaag
301 ataagttctg aacgtcgaaa agaaaagtct cgagatgcag ccagatctcg gcgaagtaaa
361 qaatctgaag ttttttatga gcttgctcat cagttgccac ttccacataa tgtgagttcg
421 catcttgata aggcctctgt gatgaggctt accatcagct atttgcgtgt gaggaaactt
481 ctggatgctg gtgatttgga tattgaagat gacatgaaag cacagatgaa ttgcttttat
541 ttgaaagcct tggatggttt tgttatggtt ctcacagatg atggtgacat gatttacatt
601 tctgataatg tgaacaaata catgggatta actcagtttg aactaactgg acacagtgtg
661 tttgatttta ctcatccatg tgaccatgag gaaatgagag aaatgcttac acacagaaat
721 ggccttgtga aaaagggtaa agaacaaaac acacagcgaa gcttttttct cagaatgaag
781 tgtaccctaa ctagccgagg aagaactatg aacataaagt ctgcaacatg gaaggtattg
841 cactgcacag gccacattca cgtatatgat accaacagta accaacctca gtgtgggtat
901 aagaaaccac ctatgacctg cttggtgctg atttgtgaac ccattcctca cccatcaaat
961 attgaaattc ctttagatag caagactttc ctcagtcgac acagcctgga tatgaaattt
1021 tottattgtg atgaaagaat taccgaattg atgggatatg agccagaaga acttttaggc
1081 cgctcaattt atgaatatta tcatgctttg gactctgatc atctgaccaa aactcatcat
1141 gatatgttta ctaaaggaca agtcaccaca ggacagtaca ggatgcttgc caaaagaggt
1201 ggatatgtct gggttgaaac tcaagcaact gtcatatata acaccaagaa ttctcaacca
1261 cagtgcattg tatgtgtgaa ttacgttgtg agtggtatta ttcagcacga cttgattttc
1321 tecetteaac aaacagaatg tgteettaaa eeggttgaat etteagatat gaaaatgaet
1381 cagctattca ccaaagttga atcagaagat acaagtagcc tctttgacaa acttaagaag
1441 gaacctgatg ctttaacttt gctggcccca gccgctggag acacaatcat atctttagat
1501 tttggcagca acgacacaga aactgatgac cagcaacttq aggaaqtacc attatataat
1561 gatgtaatgc tcccctcacc caacgaaaaa ttacaqaata taaatttqqc aatqtctcca
1621 ttacccaccg ctgaaacgcc aaagccactt cgaagtagtg ctgaccctgc actcaatcaa
1681 gaagttgcat taaaattaga accaaatcca gagtcactgg aactttcttt taccatgccc
1741 cagattcagg atcagacacc tagtccttcc gatggaagca ctagacaaag ttcacctgag
1801 cctaatagtc ccagtgaata ttgtttttat gtggatagtg atatggtcaa tgaattcaag
1861 ttggaattgg tagaaaaact ttttgctgaa gacacagaag caaagaaccc attttctact
1921 caqqacacag atttagactt ggagatgtta gctccctata tcccaatgga tgatgacttc
1981 cagttacgtt ccttcgatca gttgtcacca ttagaaagca gttccgcaag ccctgaaagc
2041 gcaagtcctc aaagcacagt tacagtattc cagcagactc aaatacaaga acctactgct
2101 aatgccacca ctaccactgc caccactgat gaattaaaaa cagtgacaaa agaccgtatg
2161 gaagacatta aaatattgat tgcatctcca tctcctaccc acatacataa agaaactact
2221 agtgccacat catcaccata tagagatact caaagtcgga cagcctcacc aaacagagca
2281 ggaaaaggag tcatagaaca gacagaaaaa tctcatccaa gaagccctaa cgtgttatct
2341 gtcgctttga gtcaaagaac tacagttcct gaggaagaac taaatccaaa gatactagct
2401 ttgcagaatg ctcagagaaa gcgaaaaatg gaacatgatg gttcactttt tcaagcagta
2461 ggaattggaa cattattaca gcagccagac gatcatgcag ctactacatc actttcttqq
2521 aaacgtgtaa aaggatgcaa atctagtgaa cagaatggaa tgqagcaaaa gacaattatt
2581 ttaataccct ctgatttagc atgtagactg ctggggcaat caatggatga aagtggatta
2641 ccacagctga ccagttatga ttgtgaagtt aatgctccta tacaaggcag cagaaaccta
2701 ctgcagggtg aagaattact cagagctttg gatcaagtta actgagcttt ttcttaattt
2761 cattectttt tttggacact ggtggctcac tacctaaagc agtctattta tattttctac
2821 atctaatttt agaagcctgg ctacaatact gcacaaactt gqttagttca atttttgatc
2881 ccctttctac ttaatttaca ttaatgctct tttttagtat gttctttaat gctggatcac
2941 agacagetea ttttctcagt tttttggtat ttaaaccatt geattgeagt ageateattt
3001 taaaaaatgc acctttttat ttatttattt ttggctaggg agtttatccc tttttcgaat
```

3061 tattttaag aagatgccaa tataatttt gtaagaaggc agtaaccttt catcatgatc 3121 ataggcagtt gaaaaatttt tacacctttt ttttcacatt ttacataaat aataatgctt 3181 tgccagcagt acgtggtagc cacaattgca caatatattt tcttaaaaaa taccagcagt 3241 tactcatgga atatattctg cgtttataaa actagtttt aagaagaaat tttttttggc 3301 ctatgaaatt gttaaacctg gaacatgaca ttgttaatca tataataatg attcttaaat 3361 gctgtatggt tattattta aatgggtaaa gccattaca taatatagaa agatatgcat 3421 atatctagaa ggtatgtggc atttattgg ataaaattct caattcagag aaatcatctg 3481 atgtttctat agtcactttg ccagctcaaa agaaaacaat accctatgta gttgtggaag 3541 tttatgctaa tattgtgtaa ctgatattaa acctaaatgt tctgcctacc ctgttggtat 3601 aaagatattt tgagcagact gtaaacaaga aaaaaaaaat catgcattct tagcaaaatt 3661 gcctagtatg ttaatttgct caaaatacaa tgtttgatt tatgcacttt gtcgctatta 3721 acatcctttt tttcatgtag atttcaataa ttgagtaatt ttagaagcat tattttagga 3781 atatatagtt gtcacagtaa atatcttgtt tttctatgt acattgtaca aattttcat 3841 tccttttgct ctttgtggtt ggaaaaaaa aaaaaaaaa aaa

Sequence I.D. No. 10

MEGAGGANDKKKISSERRKEKSRDAARSRRSKESEVFYELAHQL
PLPHNVSSHLDKASVMRLTISYLRVRKLLDAGDLDIEDDMKAQMNCFYLKALDGFVMV
LTDDGDMIYISDNVNKYMGLTQFELTGHSVFDFTHPCDHEEMREMLTHRNGLVKKGKE
QNTQRSFFLRMKCTLTSRGRTMNIKSATWKVLHCTGHIHVYDTNSNQPQCGYKKPPMT
CLVLICEPIPHPSNIEIPLDSKTFLSRHSLDMKFSYCDERITELMGYEPEELLGRSIY
EYYHALDSDHLTKTHHDMFTKGQVTTGQYRMLAKRGGYVWVETQATVIYNTKNSQPQC
IVCVNYVVSGIIQHDLIFSLQQTECVLKPVESSDMKMTQLFTKVESEDTSSLFDKLKK
EPDALTLLAPAAGDTIISLDFGSNDTETDDQQLEEVPLYNDVMLPSPNEKLQNINLAM
SPLPTAETPKPLRSSADPALNQEVALKLEPNPESLELSFTMPQIQDQTPSPSDGSTRQ
SSPEPNSPSEYCFYVDSDMVNEFKLELVEKLFAEDTEAKNPFSTQDTDLDLEMLAPYI
PMDDDFQLRSFDQLSPLESSSASPESASPQSTVTVFQQTQIQEPTANATTTTATTDEL
KTVTKDRMEDIKILIASPSPTHIHKETTSATSSPYRDTQSRTASPNRAGKGVIEQTEK
SHPRSPNVLSVALSQRTTVPEEELNPKILALQNAQRKRKMEHDGSLFQAVGIGTLLQQ
PDDHAATTSLSWKRVKGCKSSEQNGMEQKTIILIPSDLACRLLGQSMDESGLPQLTSY
DCEVNAPIQGSRNLLQGEELLRALDQVN

MOP2:

Sequence I.D. No. 2

1 gcgtctgaac gtctcaaagg gccacagcga caatgacagc tgacaaggag aagaaaagga 61 gtagctcgga gaggaggaag gagaagtccc gggatgctgc gcggtgccgg cggagcaagg 121 agacggaggt gttctatgag ctggcccatg agctgcctct gccccacagt gtgagctccc 181 atctggacaa ggcctccatc atgcgactgg caatcagctt cctgcgaaca cacaagctcc 241 totoctcagt ttgctctgaa aacgagtccg aagccgaagc tgaccagcag atggacaact 301 tgtacctgaa agccttggag ggtttcattg ccgtggtgac ccaagatggc gacatgatct 361 ttctgtcaga aaacatcagc aagttcatgg gacttacaca ggtggagcta acaggacata 421 gtatctttga cttcactcat ccctgcgacc atgaggagat tcgtgagaac ctgagtctca 481 aaaatggctc tggttttggg aaaaaaagca aagacatgtc cacagagcgg gacttcttca 541 tgaggatgaa gtgcacggtc accaacagag gccgtactgt caacctcaag tcagccacct 601 ggaaggtett geactgeacg ggecaggtga aagtetacaa caactgeect ceteacaata 661 gtctgtgtgg ctacaaggag cccctgctgt cctgcctcat catcatgtgt gaaccaatcc 721 agcacceatc ccacatggac atccccctgg atagcaagac cttcctgagc cgccacagca 781 tggacatgaa gttcacctac tgtgatgaca gaatcacaga actgattggt taccaccctg 841 aggagetget tggccgetca geetatgaat tetaceatge getagaetee gagaacatga 901 ccaagagtca ccagaacttg tgcaccaagg gtcaggtagt aagtggccag taccggatgc 961 tcgcaaagca tgggggctac gtgtggctgg agacccaggg gacggtcatc tacaaccctc 1021 gcaacctgca gccccagtgc atcatgtgtg tcaactacgt cctgagtgag attgagaaga 1081 atgacgtggt gttctccatg gaccagactg aatccctgtt caagccccac ctgatggcca 1141 tgaacagcat ctttgatagc agtggcaagg gggctgtgtc tgagaagagt aacttcctat 1201 tcaccaaget aaaggaggag ceegaggage tggeecaget ggeteecace eeaggagacg 1261 ccatcatctc tctggatttc gggaatcaga acttcgagga gtcctcagcc tatggcaagg 1321 ccatcctgcc cccgagccag ccatgggcca cggagttgag gagccacagc acccagagcg 1381 aggetgggag cetgeetgee tteacegtge eecaggeage tgeeceggge ageaceaece 1441 ccaqtqccac cagcagcagc agcagctqct ccacqcccaa tagccctqaa qactattaca 1501 catctttgga taacgacctg aagattgaag tgattgagaa gctcttcgcc atggacacag 1561 aggccaagga ccaatgcagt acccagacgg atttcaatga gctggacttg gagacactgg 1621 caccetatat ceccatqqae qqqqaaqqet tecaqetaaq ceccatetqe eecqaqqaqe 1681 ggctcttggc ggagaaccca cagtccaccc cccagcactg cttcagtgcc atgacaaaca 1741 tettecagee actggeeeet gtageeeege acagteeett ceteetggae aagttteage 1801 agcagctgga gagcaagaag acagagcccg agcgccggcc catgtcctcc atcttctttg 1861 atgccggaag caaagcatcc ctgccaccgt gctgtggcca ggccagcacc cctctctct 1921 ccatgggggg cagatecaac acccagtggc ccccagatec accattacat tttgggccca 1981 caaagtgggc cgtcggggat cagcgcacag agttcttggg agcagcgccg ttggggcccc 2041 ctgtctctcc accccatgtc tccaccttca aaacaaggtc tgcaaagggt tttggggctc 2101 gaggcccaaa cgtgctgagt ccggccatgg tagccctctc caacaagctg aagctgaagc 2161 gacagetgga gtatgaaaag caageettee aggaceegag egggggggae eeacetggtg 2221 gcagcacctc acatttgatg tggaaacgga tgaagaacct caggggtggg agctgccctt 2281 tgatgccgga caagccactg agcgcaaatg tacccaatga taagctcacc caaaactcca 2341 tgaggggcct gggccatccc ctgagacatc tgccgctqcc acaqcctcca tctqccatca 2401 gtcccgggga gaacagcaag agcaggttcc ccccacaqtg ctacgccacc cagtaccagg 2461 actacageet gtegteagee cacaaggtgt caggeatgge aageeggetg etegggeeet 2521 catttgagtc ctacctgctg cccgaactga ccagatatga ccgtgaggtg aaagtgcccg 2581 tgctgggaag ctccacgctc ctgcaaggag gggacctcct cagagccctg gaccaggcca 2641 cctgagccag gcttctacct gggcagcacc tctgccgacg ccgtcccacc agcttcactc 2701 tctccgtctg tctttgcaac taggtatttg

Sequence I.D. No. 11

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MOP3:

Sequence I.D. No. 3

1 ggagatgagc aaggaggccg tgagcctgtg ggcgctcact gtgtccctcc aaccccagt 61 ccccttgtgt gtctgcagag agatgacagg atcaggcaga agaaaacagc aatgtgtaac 121 tttgccattc atctccagag aattatgtt ttatcttttg cttttcctc cccccaggtt 181 agaatataca gaacaccaag gagggataaa aaatgcaagg gaagctcaca gtcagattga 241 aaagcggcgt cgggataaaa tgaacagttt tatagatgaa ttggcttctt tggtaccaac

301 atgcaacgca atgtccagga aattagataa acttactgtg ctaaggatgg ctgttcagca 361 catgaaaaca ttaagaggtg ccaccaatcc atacacagaa gcaaactaca aaccaacttt 421 tctatcagac gatgaattga aacacctcat tctcagggca gcagatggat ttttgtttgt 481 cgtaggatgt gaccgaggga agatactctt tgtctcagag tctgtcttca agatcctcaa 541 ctacagccag aatgatctga ttggtcagag tttgtttgac tacctgcatc ctaaagatat 601 tgccaaagtc aaggagcagc tctcctcctc tgacaccgca ccccqqqaqc qqctcataqa 661 tgcaaaaact ggacttccag ttaaaacaga tataacccct gggccatctc gattatgttc 721 tggagcacga cgttctttct tctgtaggat gaaqtgtaac aggccttcag taaaqgttga 781 agacaaggac ttcccctcta cctgctcaaa gaaaaaagca gatcgaaaaa gcttctgcac 841 aatccacagc acaggctatt tgaaaagctg gccacccaca aagatggggc tggatgaaga 901 caacgaacca gacaatgagg ggtgtaacct cagctgcctc gtcgcaattg gacgactgca 961 ttctcatgta gttccacaac cagtgaacgg ggaaatcagg gtgaaatcta tggaatatgt 1021 ttctcggcac gcgatagatg gaaagtttgt ttttgtagac cagagggcaa cagctatttt 1081 ggcatattta ccacaagaac ttctaggcac atcgtgttat gaatattttc accaagatga 1141 cataggacat cttgcagaat gtcataggca agttttacag acgagagaaa aaattacaac 1201 taattgctat aaatttaaaa tcaaagatgg ttcttttatc acactacgga gtcgatggtt 1261 cagtttcatg aacccttgga ccaaggaagt agaatatatt gtctcaacta acactgttgt 1321 tttagccaac gtcctggaag gcggggaccc aaccttccca cagctcacag catcccccca 1381 cagcatggac agcatgctgc cctctggaga aggtggccca aagaggaccc accccactgt 1441 tccagggatt ccagggggaa cccgggctgq qqcagqaaaa ataqqccqaa tqattqctqa 1501 ggaaatcatg gaaatccaca ggataagagg gtcattgcgt tctagctgtg gctccagccc 1561 attgaacatc acgagtacgc ctccccctga tgcctcttct ccaggaggca agaagatttt 1621 aaatggaggg actccagaca ttccttccag tggcctacta tcaggccagg ctcaggagaa 1681 cccaggttat ccatattctg atagttcttc tattcttggt gagaaccccc acataggtat 1741 agacatgatt gacaacgacc aaggatcaag tagtcccagt aatgatgagg cagcaatggc 1801 tgtcatcatg agcctcttgg aagcagatge tggactgggt ggccctgttg actttagtga 1861 cttgccatgg ccgctgtaaa cantacatgt tgctttggca acagcctata gtatcaaagt 1921 gcattactgg tggagtttta cagtctgtga agcttactgg ataaggagag aatagctttt 1981 atgtactgac ttcataaaag ccatctcaga gccattgata caagtcaatc ttactatatg 2041 taacttcaga caaagtggaa ctaagcctgc tccagtgttt cctcatcatt qattattggq 2101 ctagctgtgg atagcttgca ttaattgtat attttggatt ctgtttgtgt tgaatttttt 2161 aatcattgtg cacagaagca tcattggtag cttttatatg caaatggtca tttcagatgt 2221 atggtgtttt tacactacaa agaagteece catgtggata tttettatae taattgtate 2281 ataaagccgt ttattcttcc ttgtaagaat cctttactat aaatatgggt taaagtataa 2401 aaaaaaaaa aanattcqtq cqqccqctaq

Sequence I.D. No. 12

MSKEAVSLWALTVSLQPPVPLCVCREMTGSGRRKQQCVTLPFIS
RBLCFYLLLFPPPRLEYTBHQGGIKNAREAHSQIEKRRRDKMNSFIDELASLVPTCNA
MSRKLDKLTVLRMAVQHMKTLRGATNPYTEANYKPTFLSDDELKHLILRAADGFLFVV
GCDRGKILFVSESVFKILNYSQNDLIGQSLFDYLHPKDIAKVKEQLSSSDTAPRERLI
DAKTGLPVKTDITPGPSRLCSGARRSFFCRMKCNRPSVKVEDKDFPSTCSKKKADRKS
FCTIHSTGYLKSWPPTKMGLDEDNEPDNEGCNLSCLVAIGRLHSHVVPQPVNGEIRVK
SMEYVSRHAIDGKFVFVDQRATAILAYLPQELLGTSCYEYFHQDDIGHLAECHRQVLQ
TREKITTNCYKFKIKDGSFITLRSRWFSFMNPWTKEVEYIVSTNTVVLANVLEGGDPT
FPQLTASPHSMDSMLPSGEGGPKRTHPTVPGIPGGTRAGAGKIGRMIAEEIMEIHRIR
GSLRSSCGSSPLNITSTPPPDASSPGGKKILNGGTPDIPSSGLLSGQAQENPGYPYSD
SSSILGENPHIGIDMIDNDQGSSSPSNDEAAMAVIMSLLEADAGLGGPVDFSDLPWPL

MOP4:

Sequence I.D. No. 4

1 qaattccqqq ccqqaaaaac tqcataqaaa atttaatqqa tqaaqatqaq aaaqacaqaq ·61 ccaaqaqaqc ttctcqaaac aaqtctgaqa aqaaqcqtcq qqaccaqttc aatqttctca 121 teaaagaget cagttecatg etceetggea acaegeggaa aatggacaaa accaeegtgt 181 tqqaaqaqqt catcqqattt ttqcaqaaac acaatqaaqt ctcaqcqcaa acqqaaatct 241 gtgacattca gcaagactgg aagccttcat tcctcagtaa tgaagaattc acccagctga 301 tqttqqaqqc attaqatqqc ttcattatcq caqtqacaac agacqqcaqc atcatctatq 361 tetetgacag tateacgeet etcettggge atttacegte ggatgteatg gateagaatt 421 tgttaaattt cctcccagaa caagaacatt cagaagttta taaaatcctt tcttcccata 481 tgcttgtgac ggattccccc tccccagaat acttaaaatc tgacggcgat ttagaqtttt 541 attgccatct tctcaqaggc agcttgaacc caaaggaatt tccaacttat gaatacataa 601 aatttgtagg aaattttege tettacaaca atgtgeetag ecceteetgt aatggttttg 661 acaacacct ttcaagacct tgccgggtgc cactaggaaa ggaggtttgc ttcattgcca 721 ccgttcgtct ggcaacacca caattcttaa aggaaatgtg catagttgac gaacctttag 781 aggaattcac ttcaaggcat agcttggaat ggaaattttt atttctggat cacagagcac 841 ctccaatcat aggatacctg ccttttgaag tgctgggaac ctcaggctat gactactacc 901 acattgatga cctggagctc ctggccaggt gtcaccagca cctgatgcag tttggcaaag 961 ggaagtcgtg ttgctaccgg tttctgacca aaggtcagca gtggatctgg ctgcagactc 1021 actactacat cacctaccat cagtggaact ccaagcccga gttcatcgtg tgcacacact 1081 cggtggtcag ttacgcagat gtccgggtgg aaaggaggca ggagctggct ctggaagacc 1141 cqccatccqa qqccctccac tcctcaqcac taaaqqacaa qqqctcaaqc ctqqaacctc 1201 ggcagcactt taacgcactc gacgtgggtg cctcgggcct taataccagt cattcgccat 1261 eggegteete aagaagttee cacaaateet egcacaeage catgteagaa eccaeeteea 1321 ctcccaccaa gctgatggca gaggccagca ccccggcttt gccaaqatca qccaccctqc 1381 cccaagagtt acctgtcccc gggctcagcc aggcagccac catgccggcc cctctgcctt 1441 ccccatcgtc ctgcgacctc acacagcagc tcctgcctca gaccgttctg cagagcacgc 1501 ccgctcccat ggcacagttt tcggcacagt tcagcatgtt ccagaccatc aaagaccagc 1561 tagagcagcg gacgcggatc ctgcaggcca atatccggtg gcaacaggaa gagctccaca 1621 agatecagga geagetetge etggtecagg actecaaegt ceagatgtte etgeageage 1681 cagetgtate cetgagette ageageacee agegaeetga ggeteageag cagetacage 1741 aaaggtcagc tgcagtgact cagccccagc tcggggcggg cccccaactt ccagggcaga 1801 tetectetge ceaggteaca agecageace tgeteagaga ateaagtgtg atateaacee 1861 agggtccaaa gccaatgaga agctcacagc taatgcagag cagcggccgc tc

Sequence I.D. No. 13

MDEDEKDRAKRASRNKSEKKRRDQFNVLIKELSSMLPGNTRKMD
KTTVLEEVIGFLQKHNEVSAQTEICDIQQDWKPSPLSNEEFTQLMLEALDGFIIAVTT
DGSIIYVSDSITPLLGHLPSDVMDQNLLNFLPEQEHSEVYKILSSHMLVTDSPSPEYL
KSDGDLEFYCHLLRGSLNPKEFPTYEYIKFVGNFRSYNNVPSPSCNGFDNTLSRPCRV
PLGKEVCFIATVRLATPQFLKEMCIVDEPLEEFTSRHSLEWKFLFLDHRAPPIIGYLP
FEVLGTSGYDYYHIDDLELLARCHQHLMQFGKGKSCCYRFLTKGQQWIWLQTHYYITY
HQWNSKPEFIVCTHSVVSYADVRVERRQELALEDPPSEALHSSALKDKGSSLEPRQHF
NALDVGASGLNTSHSPSASSRSSHKSSHTAMSEPTSTPTKLMAEASTPALPRSATLPQ
ELPVPGLSQAATMPAPLPSPSSCDLTQQLLPQTVLQSTPAPMAQFSAQFSMFQTIKDQ
LEQRTRILQANIRWQQEELHKIQEQLCLVQDSNVQMFLQQPAVSLSFSSTQRPEAQQQ
LQQRSAAVTQPQLGAGPQLPGQISSAQVTSQHLLRESSVISTQGPKPMRSSQLMQSSG

MOP5:

Sequence I.D. No. 5

- 1 gaattcccgg agaccagcgc tgcgggccgc ggcggctggg gcgaggccag ctggggccc 61 cggctctcag ccccagagc agcacctggg aggtcacatc ttgcagtccc tggatqqctt
- 121 tgtgttcgcc ttgaaccagg aaggaaaatt cctctacatc tcagagacag tctccatcta

181 tctgggtctc tcacaggtgg agatgacggg cagcagcgtc ttcgactaca ttcaccctgg 241 ggaccactca gaggtgctgg agcaactggg gctgcggacg ccgacgcccg gcccccaac 301 cocgcottc gtotoctott cotoctott ttoctottcg ottgoagata coccogagat 361 cgaggccagc ctcaccaagg tgccccctc ctccctggtc caggagcgct ccttctttgt 421 ccgcatgaaa tccacgctca ccaagagggg gctgcacgtc aaggcctcag ggtacaaggt 481 catccacgtg actgggcgcc ttcgggccca cgccctgggc cttgtggccc tcgggcacac 541 gttgcccccg gccccctgg ctgagctgcc actccatgga cacatgatcg tcttccgtct 601 cagcctgggt ctcaccatcc ttgcttgtga gagcagagtc agcgaccaca tggacctggg 661 gccctcagag ctggtgggcc gcagctgcta ccagtttgtc cacggacaag acgccacgag 721 gatccgccag agccacgtgg acttgctgga caagggtcag gtgatgactg gttactaccg 781 ttggctgcag cgtgccgggg gcttcgtgtg gctgcagtct gtggccacag tggctgggag 841 cgggaagage cccggggage accatgtgct ttgggtcage cacgtgctca gccaagccga 901 gggtggccaa acteetttgg atgeetteca gettecagee agegtggeet gtgaggagge 961 atccagcccg gggccagagc ccacagagcc ggagcctccg acggaaggga agcaggctqc 1021 cccagcggag aacgaggccc cccagaccca gggcaaacgc atcaaagtgg agcccggccc 1081 gagggaaacc aaaggeteeg aggacagtgg cgaegaggat ceeteeagee acceggeeac 1141 accgaggece gagtteacet etgteateeg ggeagggte etgaageagg atceggtgeg 1201 gccatggggc ctggcgcctc ccggqgaccc cccqcccacc ctcctqcacq cqqqcttcct 1261 gccgccggtg gtgcggggcc tgtgcacacc cggcaccate cgctacggcc ccgcggagct 1321 gggcctggtg tacccgcacc tgcagaggct gggtccgggc cccqcgctcc cqgaggcctt 1381 ttacccgccc ctgggcctgc cctacccggg gcccgcgggc accaggctgc cgcggaaggg 1441 ggactgagga ctggcagagc tgccggcgcc ggaccctgcg acaaccgggg tcccccagga 1501 cagtaggece ggetetgece gtagecetga gaattaaaeg eeggetetee etgeaaaaaa 1561 aaaaaaaaa aaatttcctg c

Sequence I.D. No. 14

NSRRPALRAAAAGARPAGGPGSQPPEQHLGGHILQSLDGFVFAL
NQEGKFLYISETVSIYLGLSQVEMTGSSVFDYIHPGDHSEVLEQLGLRTPTPGPPTPP
SVSSSSSSSSSLADTPEIEASLTKVPPSSLVQERSFFVRMKSTLTKRGLHVKASGYKV
IHVTGRLRAHALGLVALGHTLPPAPLAELPLHGHMIVFRLSLGLTILACESRVSDHMD
LGPSELVGRSCYQFVHGQDATRIRQSHVDLLDKGQVMTGYYRWLQRAGGFVWLQSVAT
VAGSGKSPGEHHVLWVSHVLSQAEGGQTPLDAFQLPASVACEEASSPGPEPTEPEPPT
EGKQAAPAENEAPQTQGKRIKVEPGPRETKGSEDSGDEDPSSHPATPRPEFTSVIRAG
VLKQDPVRPWGLAPPGDPPTLLHAGFLPPVVRGLCTPGTIRYGPAELGLVYPHLQRL
GPGPALPEAFYPPLGLPYPGPAGTRLPRKGD

MOP6:

SEQ ID NO:6 (MOP6 cDNA)

AGCAGTGTCTTTGACTATGTCCACCCGGAGATCACGTGGAGATGGCTGAGCAGCTGGGCATGAAGCTC ${\tt CCCCTGGGGGGGTCTCCTGTCACAGGGCACTGCTGAGGACGGAGCCAGCTCAGCATCTTCCTCCTCT}$ CAGTCGGAGACCCCCGAGCCAGTGGAGTCAACCAGCCCCAGTCTGCTAACCACTGACAACACTCTTGAG CGTTCCTTTTCATCCGAATGAAATCTACTCTGACCAAACGCGGTGTGCACATCAAATCATCAGGATAT AAGGTGATTCACATAACAGGCCGGCTACGCCTGAGAGTGTCGCTGTCCCACGGGAGGACCGTCCCCAGC CAAATCATGGGTCTCGTGGTTGTTGCGCATGCCTTCCCCCCTACGATCAATGAAGTCAGAATTGAC TGCCATATGTTCGTCACTCGAGTAAATATGGACCTCAATATCATTTACTGTGAAAATAGGATTAGTGAT TATATGGATCTGACCCCTGTAGATATCGTAGGGAAGAGATGCTACCACTTCATCCATGCTGAAGACGTG GAGGGCATCAGGCACAGTCACTTGGACTTGCTGAATAAGGGTCAGTGTGTGACAAAGTACTATCGCTGG ATGCAGAAGAACGGAGGATATATTTGGATACAGTCCAGTGCCACCATAGCTATTAATGCCAAGAATGCA ANTGANAGANTATCATCTGGGTGAATTACCTTCTTAGCAATCCTGAGTACAAGGACACACCCATGGAC **ATCGCACAGCTCCCCCATCTGCCGGAGAAAACTTCCGGAATCCTCGGAGACATCCGACTCTGAGTCAGAC** TCTAAAGACACCTCAGGTATTACAGAGGACAACGAGAACTCCAAGTCCGACGAGAAGGGGAACCAGTCC GAGAACAGCGAAGACCCGGACCCGACCGGAAGAAGTCGGGCAACGCGTGTGACAACGACATGAACTGC **AACGACGCCCACAGCTCCAGTAACCCGGACAGCCGCGACAGCGACAGCTTCGAGCACTCGGAC** TTTGAGAACCCCAAGGCGGGCGAGGACGGCTTCGGTGCTCTGGGCGCGATGCAGATCAAGGTGGAGCGC TACGTGGAGAGCGAGTCGGACCTGCGGCTGCAGAACTGCGAGTCACCTCACGTCCGACAGCGCCAAGGAC CGGCAAAAGGGCGCCAGCCGCCGCCGCCCTGTCCAGCCGTCGAGCCCAGGCGGCCTGGACGCG GGCCTGGTGGAGCCCCCGCGGCTGCTGCTCCCCCAACAGTGCCTCGGTGCTCAAGATCAAGACGGAG ATCTCAGAACCCATCAATTTCGACAATGACAGCAGCATCTGGAACTACCCGCCCAACCGGGAGATCTCC **AGGAACGAGTCCCCTACAGCATGACCAAGCCCCCCAGCTCTGAGCACTTCCCGTCCCCGCAGGGCGGC** GGCGGTGGGGGTGGCGGGGGGGCTGCACGTGGCCATTCCCGACTCGGTCCTCACSCCGCCCGGC GCCGACGGC

SEQ ID NO:15 (MOP6 protein)

MAPTKPSFQQDPSRRERLQALRKEKSRDAARSRRGKENFEFYELAKLLPLPAAITSQLDKASIIRLTIS
YLKMRDFANQGDPPWNLRMEGPPPNTSVKGAQRRRSPSALAIEVFEAHLGSHILQSLDGFVFALNQEGK
FLYISETVSIYLGLSQVELTGSSVFDYVHPGDHVEMAEQLGMKLPPGRGLLSQGTAEDGASSASSSSQS
ETPEPVESTSPSLLTTDNTLERSFFIRMKSTLTKRGVHIKSSGYKVIHITGRLRLRVSLSHGRTVPSQI
MGLVVVAHALPPPTINEVRIDCHMFVTRVNMDLNIIYCENRISDYMDLTPVDIVGKRCYHFIHAEDVEG
IRHSHLDLLNKGQCVTKYYRWMQKNGGYIWIQSSATIAINAKNANEKNIIWVNYLLSNPEYKDTPMDIA
QLPHLPEKTSESSETSDSESDSKDTSGITEDNENSKSDEKGNQSENSEDPEPDRKKSGNACDNDMNCND
DGHSSSNPDSRDSDDSFEHSDFENPKAGEDGFGALGAMQIKVERYVESESDLRLQNCESLTSDSAKDSD
SAGEAGAQASSKHQKRKKRRKRQKGGSASRRRLSSASSPGGLDAGLVEPPRLLSSPNSASVLKIKTEIS
EPINFDNDSSIWNYPPNREISRNESPYSMTKPPSSEHFPSPQGGGGGGGGGGGLHVAIPDSVLTPPGAD

M P7:

Sequence I.D. No. 7 (Mouse mop7 cDNA sequence), total 2207 nucloetides):

1 agctaagtcc cggagaggac agagggcctt aggcacacaa cctaggggag aagcctggag 61 caaagcccca cagggagggc cacatggact gggaccaaga caggtcgaac accgagctqc 121 ggaaggagaa gtegegggae geggeeegea geeggegeag ceaggagaeg gaggtgetgt 181 accagetgge geacactetg ecetttgege geggegteag egegeacetg gacaaggeet 241 ccatcatgcg cctcacaatc agctacctgc gcatgcaccg cctctqcqca qcaqqqqaqt 301 ggaaccaggt ggaaaaaggg ggagagccac tggacgcctg ctacctgaag gccctggagg 361 gtttcgtcat ggtactcacc gccgagggag acatggctta cctgtcggaa aatgtcagca 421 agcacctggg cctcagtcag ctggagctca ttggacacag tatctttgat tttatccatc 481 cctgtgacca agaggaactt caagacgccc tgaccccag gccgaacctg tcaaagaaqa 541 agctggaagc cccaacagag cgccactttt ccctgcgaat gaagagcacg ctcaccaqca 601 gagggggcac gctcaacctc aaagcggcca cctggaaggt gctgcactgc tcaggacata 661 tgagggccta caagccccct gcacagactt cccctgccgg gagccctcgc tccgagcctc 721 ccctgcaatg cctggtgctt atctgtgaag ccatcccca cccagccagt ctggagcccc 781 cgctgggccg aggggccttt ctcagtcgcc acagcctgga catgaagttc acatactgcg 841 acgagaggat tgcagaagtt gctggctaca gtcctgatga cctgattggc tgttctgcct 901 atgaatacat ccacgetttg gactetgatg eggteageag gageateeac aetttgttga 961 gcaagggcca ggcagtaacg gggcagtatc gcttcctggc ccggactgga ggctatctgt 1021 ggactcagac tcaggctaca gtggtgtcag gggggcgggg cccccagtcg gaaagtatca 1081 tetgegteca ettectgate ageegtgtag aagagacegg agtggtgetg tetetggaae 1141 aaacggagca acatactcgc agaccccctc ggctgagtgc ctcctcgcag aagggtatcc 1201 ctggcaacag tgtagactct cctgctccgc ggatcctggc cttcctgcac cctccqqccc 1261 tgagtgaggc ctccctggct gctgaccctc gccgtttctg tagtccagac ctgcgccgcc 1321 tcatggcacc catcctggat ggacctcccc cagctgccac gcccagcacc ccacaagcta 1381 caeggagace ecaaagteet etteeggetg ateteceaga taagttggea gtgggettgg 1441 aaaatgcgca cagactctcc actgcccaga aaaacaagac cgtggagaca gatctagata 1501 tagetcagga etetgacact etggacttgg agatgetgge eccetacate tecatggatg 1561 atgacttcca gctcaactcc agtgagcaat tgcccaaagt ccaccgcaga cctcccaggg 1621 tggcccgcag gccccgtgct cggagcttcc atggcctgtc gcctcctatc cctgagccct 1681 ccctactgcc ccgctggggg agtgatccac gactgaactg ttccaqtcct tccaqqqqcq 1741 ategececae agecteett atgeettgaa eteggaagag ggeettggee eagageteag 1801 aggacaaagg gttggagctg ctggaaatta agcctcccaa gcggtcccca agactagaac 1861 ctggaagett cctgctgcct ccgctcagcc tgagtttcct tctgcaaggt cgacaactcc

1921 tgggaaacca gcaggatece agagecece tegtgeatte teatgageee ttgggeetag
1981 eteeeteget getetete tgeeageatg aggaaactgt ceageecagg aaccaettee
2041 egeeageage aggettggge cagacecaet gagteageet teetetaage eetettete
2101 tateeeagaa aggaeteage cacaeteeae accageagee tacaeceagg atggggeete
2161 teteetetga gtgtgeeeee eeeeageeag eeacagteet aceteag

Sequence I.D. No. 16 (Mouse mop7 protein sequence), total 662 amino acids)

MDWDQDRSNT ELRKEKSRDA ARSRRSQETE VLYQLAHTLP FARGVSAHLD KASIMRLTIS YLRMHRLCAA GEWNQVEKGG EPLDACYLKA LEGFVMVLTA EGDMAYLSEN VSKHLGLSQL ELIGHSIFDF IHPCDQEELQ DALTPRPNLS KKKLEAPTER HFSLRMKSTL TSRGRTLNLK AATWKVLHCS GHMRAYKPPA QTSPAGSPRS EPPLQCLVLI CEAIPHPASL EPPLGRGAFL SRHSLDMKFT YCDERIAEVA GYSPDDLIGC SAYEYIHALD SDAVSRSIHT LLSKGQAVTG QYRFLARTGG YLWTQTQATV VSGGRGPQSE SIICVHFLIS RVEETGVVLS LEQTEQHTRR PPRLSASSQK GIPGNSVDSP APRILAFLHP PALSEASLAA DPRRFCSPDL RRLMAPILDG PPPAATPSTP QATRRPQSPL PADLPDKLAV GLENAHRLST AQKNKTVETD LDIAQDSDTL DLEMLAPYIS MDDDFQLNSS EQLPKVHRRP PRVARRPRAR SFHGLSPPIP EPSLLPRWGS DPRLNCSSPS RGDRPTASLM PGTRKRALAQ SSEDKGLELL EIKPPKRSPR LEPGSFLLPP LSLSFLLQGR QLLGNQQDPR APLVHSHEPL GLAPSLLSLC QHEETVQPRN HFPPAAGLGQ TH

MOP8:

Sequence I.D. No. 8

1 gtggtcgage egegegeagg gtgegetegt ttgaactge gtgacacega gggttgggga 61 etegaacece egettegeag eteaggagee tgaggteega aagettegtt eeagageeea 121 geatgaatgg atacgeggaa ttteegeeea geeceagtaa eeceaceaag gageeegtgg 181 ageeceagee eageeaggte eeactgeagg aagatgtgga eatgageagt ggeteeagtg 241 gacatgagae eaacgaaaae tgeteeacgg ggegggaete geagggeagt gaetgtgaeg 301 acagtgggaa ggagetgggg atgetggtgg ageeacegga tgeeegeeag agteeagata WO 99/28464 PCT/US98/25314

361 cctttagcct gatgatggca aaatctgaac acaacccatc tacaagtggc tgcagtagcg 421 accagtette gaaagtggac acacacaaag aactgataaa aacactaaag gagetgaagg 481 tecacetece tgcagacaag aaggecaagg gcaaggecag tacgetqgec acettqaagt 541 acgccctcag gagcgtgaag caggtgaaag ccaatgaaga gtattaccag ctgctgatgt 601 ccagcgaggg tcacccctgt ggagcagacg tgccctccta caccgtggag gagatggaga 661 gegttacete tgagcacatt gtgaagaatg cegatatgtt tgeggtggee gtgteeetgg 721 tgtctgggaa gatcctgtac atctctgacc aggttgcatc catatttcac tgtaaaagag 781 atgccttcag cgatgccaag tttgtggagt tcctggcgcc tcacgatgtg ggcqtqttcc 841 acagtttcac ctccccgtac aagcttccct tgtggagcat gtgcaqtqqa qcagattctt 901 ttactcaaga atgcatggag gagaaatctt tcttttgccg tgtcagtgtc cggaaaagcc 961 acgagaatga aatccgctac caccccttcc gcatgacgcc ctacctqqtc aaqqtqcqqq 1021 accaacaagg tgctgagagt cagetttgct geettetget ggcagagaga qtgcaetetg 1081 gttatgaagc ccctagaatt cctcctgaaa agagaatttt tacaaccacc catacaccaa 1141 attgtttgtt ccaggatgtg gatgaaaggg cggtccctct cctgggctac ctacctcagg 1201 acctgattga aaccccagtg ctcgtgcagc tccaccctag tgacaggccc ttqatqctqq 1261 ccatccacaa aaagatcctg cagtcaggcg ggcagccttt cgactattct cccattcggt 1321 ttcgcgcccg gaacggagag tacatcacgt tggacaccag ctggtccagc ttcatcaacc 1381 catggagcag gaaaatctcc ttcatcattg ggaggcacaa agtcagggtg ggccctttga 1441 atgaggacgt gtttgcagcc cacccctgca cagaggagaa ggccctgcac cccagcattc 1501 aggageteac agageagate caceggetee tgetgeagee egteeceac ageggeteca 1561 gtggctacgg gagtctgggc agcaacgggt cccacgagca ccttatgagc cagacctcct 1621 ccagcgacag caacggccat gaggactcac gccggaggag agccgaaatt tqtaaaaatq 1681 gtaacaagac caaaaataga agtcattatt ctcatgaatc tggagaacaa aagaaaaaat 1741 ccgttacaga aatgcaaact aatcccccag ctgagaagaa agctqtccct gccatqqaaa 1801 aggacagect gggggteage tteccegagg agttggeetg caagaaccag eccacetget 1861 cctaccagca gatcagctgc ttggacagcg tcatcaggta cttggagagc tgcaatgagg 1921 ctgccaccct gaagaggaaa tgcgagttcc cagcaaacgt cccagcgcta aggtccaqtq 1981 ataageggaa ggccacagte ageccaggge cacacgetgg agaggcagag ecgeceteca 2041 gggtgaacag ccgcacggga gtaggtacgc acctgacctc gctggcactg ccgggcaagg 2101 cagagagtgt ggcgtcgctc accagccagt gcagctacag cagcaccatc gtccatqtqq 2161 gagacaagaa gccgcagccg gagttagaga tggtggaaga tgctgcgagt gggccagaat 2221 ccctggactg cctggcgggc cctgccctgg cctgtggtct cagccaagag aaggagccct 2281 tcaagaagct gggcctcacc aaggaggtac tcgctgcaca cacacagaag gaggagcaga 2341 getteetgea gaagtteaaa gaaataagaa aacteageat ttteeagtee cactgeeatt 2401 actacttgca agaaagatcc aaggggcagc caagtgaacg aactgcccct ggactaagaa 2461 atacttccgg aatagattca ccttggaaaa aaacaggaaa gaacagaaaa ttgaagtcca 2521 agegggteaa acctegagae teatetgaga geaeeggate tggggggeee gtgteegee 2581 ggccccgct ggtgggcttg aacgccacag cctggtcacc ctcagacacg tcccagtcca

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Sequence I.D. No. 17

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KPQPELEMVEDAASGPESLDCLAGPALACGLSQEKEPFKKLGLTKEVLAAHTQKEEQS
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MGRASPPLFQSRSSSPLQLNLLQLEEAPEGGTGAMGTTGATETAAVGADCKPGTSRDQ
QPKAPLTRDEPSDTQNSDALSTSSGLLNLLLNEDLCSASGSAASESLGSGSLGCDASP
SGAGSSDTSHTSKYFGSIDSSENNHKAKMNTGMEESEHFIKCVLQDPIWLLMADADSS
VMMTYQLPSRNLEAVLKEDREKLKLLQKLQPGSRRVRSRSCARSTSGCRRAACPQPST
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MOP9:

SEQ ID NO:9 (MOP9 cDNA)

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L

TGATGGTGCACAGTTGGATTTCGATGCCCTATGTGACAATGATGACACAGCCATGGCTGCATTTATGAATTAC
TTAGAAGCAGAGGGGGGCCTGGGAGACCCTGGGGACCTTCAGTGACATCCAGTGGACCCTCTAG

SEQ ID NO:18 (MOP9 protein)

MGSFSSHMTEFPRKRKGSDSDPSQSGIMTEKVVEKLSQNPLTYLLSTRIEISASSGSREAHSQTEKRRRDKMN
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